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NEWS 8 Mar 22 TRCTHERMO no longer available
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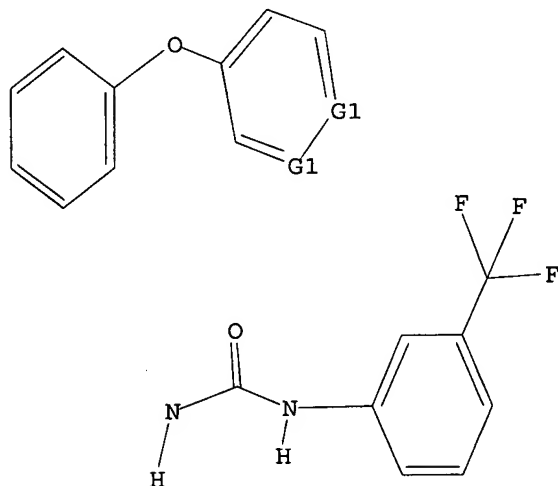
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<http://www.cas.org/ONLINE/STN/STNOTES/stnotes27.pdf>

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100.0% PROCESSED 36 ITERATIONS 13 ANSWERS
SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**
BATCH **COMPLETE**
PROJECTED ITERATIONS: 361 TO 1079
PROJECTED ANSWERS: 44 TO 476

L2 13 SEA SSS SAM L1

=> s l1 ful

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FULL SCREEN SEARCH COMPLETED - 825 TO ITERATE

100.0% PROCESSED 825 ITERATIONS 365 ANSWERS
SEARCH TIME: 00.00.02

L3 365 SEA SSS FUL L1

=> file uspatall

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FULL ESTIMATED COST	140.28	140.49

FILE 'USPATFULL' ENTERED AT 16:56:20 ON 15 JUL 2002
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FILE 'USPAT2' ENTERED AT 16:56:20 ON 15 JUL 2002
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=> s l3

L4 17 L3

=> d abs bib fhitr 1-17

L4 ANSWER 1 OF 17 USPATFULL

AB This invention relates to the use of a group of heteroaryl ureas containing nitrogen in treating p38 mediated diseases, and pharmaceutical compositions for use in such therapy.

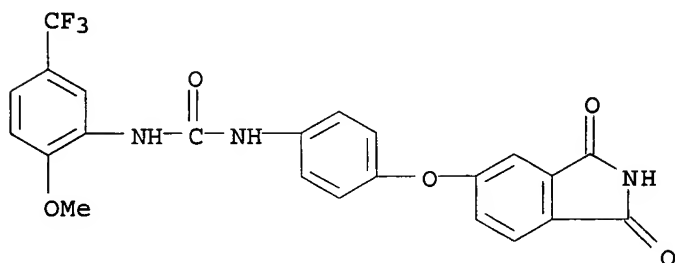
CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AN 2002:126779 USPATFULL

TI Heteroaryl ureas containing nitrogen hetero-atoms as p38 kinase inhibitors

IN Dumas, Jacques, Orange, CT, UNITED STATES
Riedl, Bernd, Wuppertal, GERMANY, FEDERAL REPUBLIC OF
Khire, Uday, Hamden, CT, UNITED STATES
Sibley, Robert N., North Haven, CT, UNITED STATES
Hatoum-Mokdad, Holia, Hamden, CT, UNITED STATES
Monahan, Mary-Katherine, Hamden, CT, UNITED STATES
Gunn, David E., Hamden, CT, UNITED STATES
Lowinger, Timotthy B., Nishinomiya City, JAPAN
Scott, William J., Guilford, CT, UNITED STATES

Smith, Roger A., Madison, CT, UNITED STATES
Wood, Jill E., Hamden, CT, UNITED STATES
PA BAYER CORPORATION (U.S. corporation)
PI US 2002065296 A1 20020530
AI US 2001-838286 A1 20010420 (9)
RLI Continuation-in-part of Ser. No. US 2001-778039, filed on 7 Feb 2001,
PENDING Continuation-in-part of Ser. No. US 1999-425229, filed on 22 Oct
1999, PENDING Continuation of Ser. No. US 1999-257265, filed on 25 Feb
1999, ABANDONED
PRAI US 1999-115878P 19990113 (60)
DT Utility
FS APPLICATION
LREP MILLEN, WHITE, ZELANO & BRANIGAN, P.C., 2200 CLARENDON BLVD., SUITE
1400, ARLINGTON, VA, 22201
CLMN Number of Claims: 39
ECL Exemplary Claim: 1
DRWN No Drawings
LN.CNT 2826
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
IT 284461-54-7P, N-[2-Methoxy-5-(trifluoromethyl)phenyl]-N'-[4-(1,3-
dioxoisindolin-5-yloxy)phenyl]urea
(prepn. of heteroaryl ureas contg. nitrogen hetero-atoms as p38 kinase
inhibitors)
RN 284461-54-7 USPATFULL
CN Urea, N-[4-[(2,3-dihydro-1,3-dioxo-1H-isindol-5-yl)oxy]phenyl]-N'-[2-
methoxy-5-(trifluoromethyl)phenyl]- (9CI) (CA INDEX NAME)

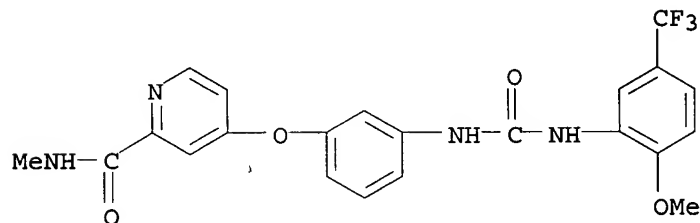


L4 ANSWER 2 OF 17 USPATFULL
AB This invention relates to the use of a group of aryl ureas in treating
raf mediated diseases, and pharmaceutical compositions for use in such
therapy.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AN 2002:78859 USPATFULL
TI Omega-carboxyaryl substituted diphenyl ureas as raf kinase inhibitors
IN Uday, Khire, Hamden, CT, UNITED STATES
Dumas, Jacques, Orange, CT, UNITED STATES
Riedl, Bernd, Wuppertal, GERMANY, FEDERAL REPUBLIC OF
Lowinger, Timothy B., Nishinomiya City, JAPAN
Scott, William J., Guilford, CT, UNITED STATES
Smith, Roger A., Madison, CT, UNITED STATES
Wood, Jill E., Hamden, CT, UNITED STATES
Monahan, Mary-Katherine, Hamden, CT, UNITED STATES
Natero, Reina, Hamden, CT, UNITED STATES
Joel, Renick, Milford, CT, UNITED STATES

Sibley, Robert N., North Haven, CT, UNITED STATES
PA BAYER CORPORATION, Pittsburgh, PA, 15205 (U.S. corporation)
PI US 2002042517 A1 20020411
AI US 2001-948915 A1 20010910 (9)
RLI Continuation of Ser. No. US 1999-425228, filed on 22 Oct 1999, ABANDONED
Continuation-in-part of Ser. No. US 1999-257266, filed on 25 Feb 1999,
ABANDONED
PRAI US 1999-115877P 19990113 (60)
DT Utility
FS APPLICATION
LREP MILLEN, WHITE, ZELANO & BRANIGAN, P.C., 2200 CLARENDON BLVD., SUITE
1400, ARLINGTON, VA, 22201
CLMN Number of Claims: 67
ECL Exemplary Claim: 1
DRWN No Drawings
LN.CNT 3675
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
IT 284461-42-3P
(prepn. of .omega.-carboxy(hetero)aryl substituted di-Ph urea raf
kinase inhibitors by reacting arylisocyanates with arylamines)
RN 284461-42-3 USPATFULL
CN 2-Pyridinecarboxamide, 4-[3-[[[2-methoxy-5-(trifluoromethyl)phenyl]amino]
carbonyl]amino]phenoxy]-N-methyl- (9CI) (CA INDEX NAME)

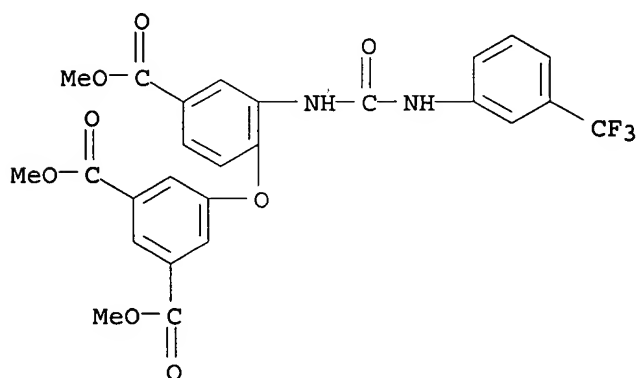


L4 ANSWER 3 OF 17 USPATFULL
AB Chemical structures have been identified which allosterically modify
pyruvate kinase and inhibit enzymatic activity. These compounds can be
used as pharmaceuticals in the treatment of a wide variety of diseases
and disorders where influencing metabolic processes is beneficial, such
as the glycolytic pathway, all pathways which use ATP as an energy
source, and all pathways which involve 2,3-diphosphoglycerate related to
the delivery of oxygen by modifying hemoglobin's oxygen affinity,
treatments of tumor and cancer and Alzheimer's disease (AD).

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AN 2001:218507 USPATFULL
TI Allosteric inhibitors of pyruvate kinase
IN Abraham, Donald J., Midlothian, VA, United States
Wang, Changging, Richmond, CA, United States
Danso-Danquah, Richmond, VA, United States
Burnett, James C., Ashland, VA, United States
Joshi, Gajanan S., Glen Allen, VA, United States
Hoffman, Steven J., Carlisle, MA, United States
PI US 2001046997 A1 20011129
AI US 2001-799873 A1 20010307 (9)
RLI Continuation-in-part of Ser. No. US 1998-46643, filed on 24 Mar 1998,
GRANTED, Pat. No. US 6214879

DT Utility
FS APPLICATION
LREP McGuire Woods, LLP, Suite 1800, 1750 Tysons Boulevard, Tysons Corner,
McLean, VA, 22102
CLMN Number of Claims: 24
ECL Exemplary Claim: 1
DRWN 7 Drawing Page(s)
LN.CNT 688
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
IT 289060-07-7
(pyruvate kinase allosteric inhibitors for therapeutic use)
RN 289060-07-7 USPATFULL
CN 1,3-Benzenedicarboxylic acid, 5-[4-(methoxycarbonyl)-2-[[[3-(trifluoromethyl)phenyl]amino]carbonyl]amino]phenoxy]-, dimethyl ester
(9CI) (CA INDEX NAME)

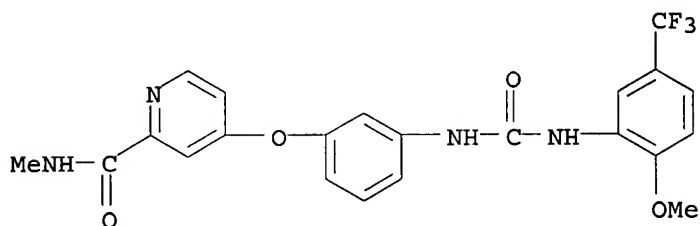


L4 ANSWER 4 OF 17 USPATFULL
AB This invention relates to the use of a group of aryl ureas in treating raf mediated diseases, and pharmaceutical compositions for use in such therapy.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AN 2001:188813 USPATFULL
TI Omega-carboxyaryl substituted diphenyl ureas as raf kinase inhibitors
IN Riedl, Bernd, Wuppertal, Germany, Federal Republic of
Dumas, Jacques, Orange, CT, United States
Khire, Uday, Hamden, CT, United States
Lowinger, Timothy P., Nashnomya City, Japan
Scott, William J., Guilford, CT, United States
Smith, Roger A., Madison, CT, United States
Wood, Jill E., Hamden, CT, United States
Monahan, Mary-Katherine, Hamden, CT, United States
Natero, Rena, Hamden, CT, United States
Renick, Joel, Milford, CT, United States
Sibley, Robert N., North Haven, CT, United States
PI US 2001034447 A1 20011025
AI US 2001-773604 A1 20010202 (9)
RLI Continuation of Ser. No. US 1999-425228, filed on 22 Oct 1999, PENDING
Continuation-in-part of Ser. No. US 1999-257266, filed on 25 Feb 1999,
ABANDONED
PRAI US 1999-115877P 19990113 (60)
DT Utility

FS APPLICATION
LREP MILLEN, WHITE, ZELANO & BRANIGAN, P.C., 2200 CLARENDON BLVD., SUITE
1400, ARLINGTON, VA, 22201
CLMN Number of Claims: 67
ECL Exemplary Claim: 1
DRWN No Drawings
LN.CNT 3666
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
IT 284461-42-3P
(prepn. of .omega.-carboxy(hetero)aryl substituted di-Ph urea raf
kinase inhibitors by reacting arylisocyanates with arylamines)
RN 284461-42-3 USPATFULL
CN 2-Pyridinecarboxamide, 4-[3-[[[2-methoxy-5-(trifluoromethyl)phenyl]amino]
carbonyl]amino]phenoxy]-N-methyl- (9CI) (CA INDEX NAME)



L4 ANSWER 5 OF 17 USPATFULL
AB This invention relates to the use of a group of aryl ureas in treating
raf mediated diseases, and pharmaceutical compositions for use in such
therapy.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AN 2001:171152 USPATFULL
TI Omega-carboxyaryl substituted disphenyl ureas as raf kinase inhibitors
IN Riedl, Bernd, Wuppertal, Germany, Federal Republic of
Dumas, Jacques, Orange, CT, United States
Khire, Uday, Hamden, CT, United States
Lowinger, Timothy B., Nishinomiya City, Japan
Scott, William J., Guilford, CT, United States
Smith, Roger A., Madison, CT, United States
Wood, Jill E., Hamden, CT, United States
Monahan, Mary-Katherine, Hamden, CT, United States
Natero, Reina, Hamden, CT, United States
Renick, Joel, Milford, CT, United States
Sibley, Robert N., Noth Haven, CT, United States
PI US 2001027202 A1 20011004
AI US 2001-773658 A1 20010202 (9)
RLI Continuation of Ser. No. US 1999-425228, filed on 22 Oct 1999, PENDING
Continuation-in-part of Ser. No. US 1999-257266, filed on 25 Feb 1999,
ABANDONED
PRAI US 1999-115877P 19990113 (60)
DT Utility
FS APPLICATION
LREP MILLEN, WHITE, ZELANO & BRANIGAN, P.C., Arlington Courthouse Plaza I,
Suite 1400, 2200 Clarendon Boulevard, Arlington, VA, 22201
CLMN Number of Claims: 67
ECL Exemplary Claim: 1
DRWN No Drawings

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LN.CNT 3656

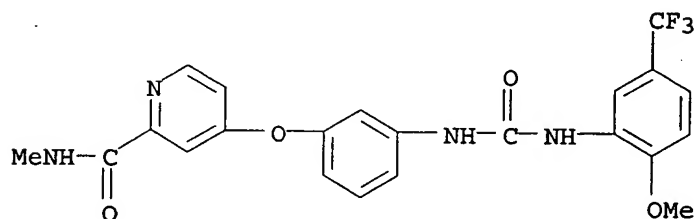
CAS INDEXING IS AVAILABLE FOR THIS PATENT.

IT 284461-42-3P

(prepn. of .omega.-carboxy(hetero)aryl substituted di-Ph urea raf kinase inhibitors by reacting arylisocyanates with arylamines)

RN 284461-42-3 USPATFULL

CN 2-Pyridinecarboxamide, 4-[3-[[[2-methoxy-5-(trifluoromethyl)phenyl]amino]carbonyl]amino]phenoxy]-N-methyl- (9CI) (CA INDEX NAME)



L4 ANSWER 6 OF 17 USPATFULL

AB This invention relates to the use of a group of aryl ureas in treating raf mediated diseases, and pharmaceutical compositions for use in such therapy.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AN 2001:139616 USPATFULL

TI Omega-carboxyaryl substituted diphenyl ureas as raf kinase inhibitors

IN Riedl, Bernd, Wuppertal, Germany, Federal Republic of

Dumas, Jacques, Orange, CT, United States

Khire, Uday, Hamden, CT, United States

Lowinger, Timothy B., Nashnomya City, Japan

Scott, William J., Guilford, CT, United States

Smith, Roger A., Madison, CT, United States

Wood, Jill E., Hamden, CT, United States

Monahan, Mary-Katherine, Hamden, CT, United States

Natéro, Rena, Hamden, CT, United States

Renick, Joel, Milford, CT, United States

Sibley, Robert N., North Haven, CT, United States

PI US 2001016659 A1 20010823

AI US 2001-773672 A1 20010202 (9)

RLI Continuation of Ser. No. US 1999-425228, filed on 22 Oct 1999, PENDING
Continuation-in-part of Ser. No. US 1999-257266, filed on 25 Feb 1999,
ABANDONED

PRAI US 1999-115877P 19990113 (60)

DT Utility

FS APPLICATION

LREP MILLEN, WHITE, ZELANO & BRANIGAN, P.C., 2200 CLARENDON BLVD., SUITE
1400, ARLINGTON, VA, 22201

CLMN Number of Claims: 67

ECL Exemplary Claim: 1

DRWN No Drawings

LN.CNT 3652

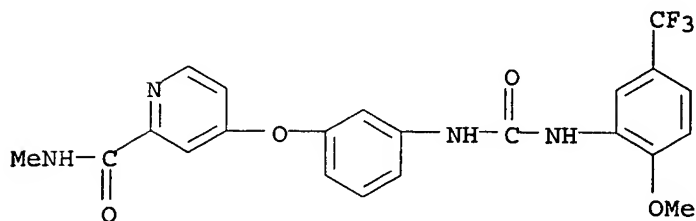
CAS INDEXING IS AVAILABLE FOR THIS PATENT.

IT 284461-42-3P

(prepn. of .omega.-carboxy(hetero)aryl substituted di-Ph urea raf kinase inhibitors by reacting arylisocyanates with arylamines)

RN 284461-42-3 USPATFULL

CN 2-Pyridinecarboxamide, 4-[3-[[[2-methoxy-5-(trifluoromethyl)phenyl]amino]carbonyl]amino]phenoxy]-N-methyl- (9CI) (CA INDEX NAME)



L4 ANSWER 7 OF 17 USPATFULL

AB This invention relates to the use of a group of aryl ureas in treating raf mediated diseases, and pharmaceutical compositions for use in such therapy.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AN 2001:123628 USPATFULL

TI omega-carboxyyaryl substituted diphenyl ureas as raf kinase inhibitors

IN Riedl, Bernd, Wuppertal, Germany, Federal Republic of

Dumas, Jacques, Orange, CT, United States

Khire, Uday, Hamden, CT, United States

Lowinger, Timothy B., Nishinomiya City, Japan

Scott, William J., Guilford, CT, United States

Smith, Roger A., Madison, CT, United States

Wood, Jill E., Hamden, CT, United States

Monahan, Mary-Katherine, Hamden, CT, United States

Natero, Reina, Hamden, CT, United States

Renick, Joel, Milford, CT, United States

Sibley, Robert N., North Haven, CT, United States

PI US 2001011136 A1 20010802

AI US 2001-773675 A1 20010202 (9)

RLI Continuation of Ser. No. US 1999-425228, filed on 22 Oct 1999, PENDING
Continuation-in-part of Ser. No. US 1999-257266, filed on 25 Feb 1999,
ABANDONED

PRAI US 1999-115877P 19990113 (60)

DT Utility

FS APPLICATION

LREP MILLEN, WHITE, ZELANO & BRANIGAN, P.C., Suite 1400, 2200 Clarendon
Blvd., Arlington, VA, 22201

CLMN Number of Claims: 67

ECL Exemplary Claim: 1

DRWN No Drawings

LN.CNT 3646

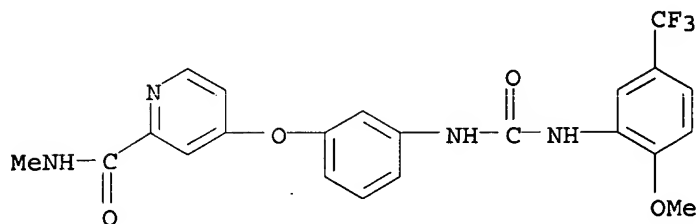
CAS INDEXING IS AVAILABLE FOR THIS PATENT.

IT 284461-42-3P

(prepn. of .omega.-carboxy(hetero)aryl substituted di-Ph urea raf
kinase inhibitors by reacting arylisocyanates with arylamines)

RN 284461-42-3 USPATFULL

CN 2-Pyridinecarboxamide, 4-[3-[[[2-methoxy-5-(trifluoromethyl)phenyl]amino]carbonyl]amino]phenoxy]-N-methyl- (9CI) (CA INDEX NAME)



L4 ANSWER 8 OF 17 USPATFULL

AB This invention relates to the use of a group of aryl ureas in treating raf mediated diseases, and pharmaceutical compositions for use in such therapy.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AN 2001:123627 USPATFULL

TI Omega-carboxyaryl substituted diphenyl ureas as raf kinase inhibitors

IN Riedl, Bernd, Wuppertal, Germany, Federal Republic of

Dumas, Jacques, Orange, CT, United States

Khire, Uday, Hamden, CT, United States

Lowinger, Timothy B., Nishinomiya City, Japan

Scott, William J., Guilford, CT, United States

Smith, Roger A., Madison, CT, United States

Wood, Jill E., Hamden, CT, United States

Monahan, Mary-Katherine, Hamden, CT, United States

Natero, Reina, Hamden, CT, United States

Renick, Joel, Milford, CT, United States

Sibley, Robert N., North Haven, CT, United States

PI US 2001011135 A1 20010802

AI US 2001-773659 A1 20010202 (9)

RLI Continuation of Ser. No. US 1999-425228, filed on 22 Oct 1999, PENDING
Continuation-in-part of Ser. No. US 1999-257266, filed on 25 Feb 1999,
ABANDONED

PRAI US 1999-115877P 19990113 (60)

DT Utility

FS APPLICATION

LREP MILLEN, WHITE, ZELANO & BRANIGAN, P.C., Suite 1400, Arlington Courthouse
Plaza 1, Arlington, VA, 22201

CLMN Number of Claims: 67

ECL Exemplary Claim: 1

DRWN No Drawings

LN.CNT 3686

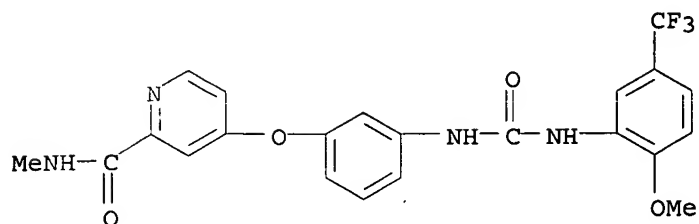
CAS INDEXING IS AVAILABLE FOR THIS PATENT.

IT 284461-42-3P

(prepn. of .omega.-carboxy(hetero)aryl substituted di-Ph urea raf
kinase inhibitors by reacting arylisocyanates with arylamines)

RN 284461-42-3 USPATFULL

CN 2-Pyridinecarboxamide, 4-[3-[[[2-methoxy-5-(trifluoromethyl)phenyl]amino]
carbonyl]amino]phenoxy]-N-methyl- (9CI) (CA INDEX NAME)



L4 ANSWER 9 OF 17 USPATFULL

AB The present invention relates to novel quinoline derivatives and quinazoline derivatives represented by the following formula (I):
 ##STR1## [wherein R.sub.1 and R.sub.2 are each independently H or C.sub.1 -C.sub.4 -alkyl, or R.sub.1 and R.sub.2 together form C.sub.1 -C.sub.3 -alkylene, X is O, S or CH.sub.2, W is CH or N, and Q is a substituted aryl group or substituted heteroaryl group] and their pharmaceutically acceptable salts, having platelet-derived growth factor receptor autophosphorylation inhibitory activity, to pharmaceutical compositions containing these compounds, and to methods for the treatment of diseases associated with abnormal cell growth such as tumors.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AN 2000:150184 USPATFULL

TI Quinoline and quinazoline derivatives inhibiting platelet-derived growth factor receptor autophosphorylation and pharmaceutical compositions containing the same

IN Kubo, Kazuo, Takasaki, Japan
 Ohyama, Shinichi, Takasaki, Japan
 Shimizu, Toshiyuki, Takasaki, Japan
 Nishitoba, Tsuyoshi, Takasaki, Japan
 Kato, Shinichiro, Takasaki, Japan
 Murooka, Hideko, Takasaki, Japan
 Kobayashi, Yoshiko, Takasaki, Japan

PA Kirin Beer Kabushiki Kaisha, Tokyo-to, Japan (non-U.S. corporation)

PI US 6143764 20001107

WO 9717329 19970515

AI US 1998-68660 19980506 (9)

WO 1996-JP3229 19961105

19980506 PCT 371 date

19980506 PCT 102(e) date

PRAI JP 1995-313555 19951107

JP 1996-62121 19960223

DT Utility

FS Granted

EXNAM Primary Examiner: Seaman, D. Margaret

LREP Foley & Lardner

CLMN Number of Claims: 52

ECL Exemplary Claim: 1

DRWN No Drawings

LN.CNT 5569

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

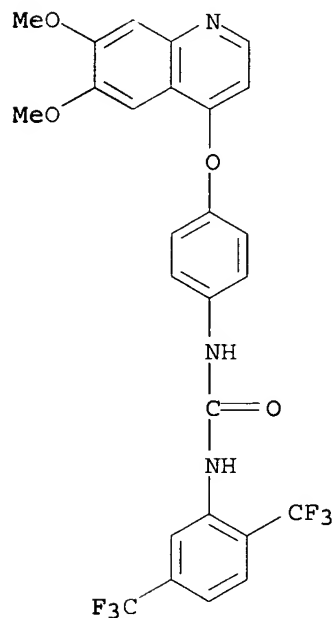
IT 190727-78-7P

(prepn. of quinoline and quinazoline derivs. inhibiting platelet-derived growth factor receptor autophosphorylation)

RN 190727-78-7 USPATFULL

CN Urea, N-[2,5-bis(trifluoromethyl)phenyl]-N'-[4-[(6,7-dimethoxy-4-

quinolinyl)oxy]phenyl]- (9CI) (CA INDEX NAME)



L4 ANSWER 10 OF 17 USPATFULL

AB This invention relates to the novel pharmaceutical compositions of Formulas (I) and (II) each of which comprises a compound of Formula (I) or (II) and a pharmaceutically acceptable diluent or carrier. This invention also relates to a method of treating or reducing inflammation in a mammal in need thereof, which comprises administering to said mammal an effective amount of a compound or composition of Formula (I) or (II).

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AN 1999:67289 USPATFULL

TI Anti-inflammatory compounds

IN Dixon, James Scott, Malvern, PA, United States

Hall, Ralph Floyd, Villanova, PA, United States

Marshall, Lisa Ann, Wyndmoor, PA, United States

Chilton, III, Floyd H., Pilot Mountain, NC, United States

Mayer, Ruth Judik, Wayne, PA, United States

Winkler, James David, Fort Washington, PA, United States

PA SmithKline Beecham Corporation, Philadelphia, PA, United States (U.S. corporation)

The Johns Hopkins University, Baltimore, MD, United States (U.S. corporation)

PI US 5912270 19990615

WO 9533712 19951214

AI US 1996-737650 19961122 (8)

WO 1995-US6677 19950602

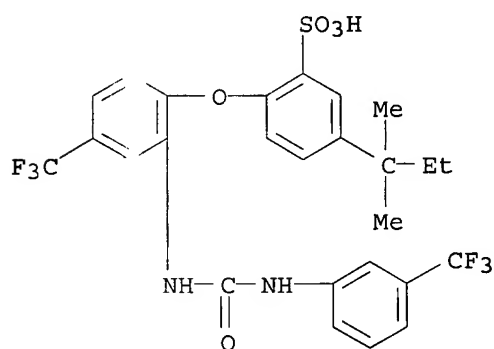
19961122 PCT 371 date

19961122 PCT 102(e) date

RLI Continuation-in-part of Ser. No. US 1994-252716, filed on 2 Jun 1994, now patented, Pat. No. US 5470882

DT Utility

FS Granted
EXNAM Primary Examiner: Gerstl, Robert
LREP Dinner, Dara L., Venetianer, Stephen, Kinzig, Charles
CLMN Number of Claims: 15
ECL Exemplary Claim: 1
DRWN No Drawings
LN.CNT 1767
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
IT 447-64-3P
 (prepn. of antiinflammatory ureidophenoxybenzenesulfonates)
RN 447-64-3 USPATFULL
CN Benzenesulfonic acid, 5-(1,1-dimethylpropyl)-2-[4-(trifluoromethyl)-2-
 [[[3-(trifluoromethyl)phenyl]amino]carbonyl]amino]phenoxy] - (9CI) (CA
 INDEX NAME)



L4 ANSWER 11 OF 17 USPATFULL
AB This invention relates to the novel pharmaceutical compositions of
 Formulas (I) and (II) each of which comprises a compound of Formula (I)
 or (II) and a pharmaceutically acceptable diluent or carrier.

 This invention also relates to a method of treating or reducing
 inflammation in a mammal in need thereof, which comprises administering
 to said mammal an effective amount of a compound or composition of
 Formula (I) or (II).

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AN 5:105872 USPATFULL
TI Anti-inflammatory compounds
IN Dixon, James S., Malvern, PA, United States
 Hall, Ralph F., Villanova, PA, United States
 Marshall, Lisa A., Wyndmoor, PA, United States
 Chilton, III, Floyd H., Pilot Mountain, NC, United States
 Payer, Ruth J., Wayne, PA, United States
 Winkler, James D., Fort Washington, PA, United States
PA SmithKline Beecham Corp., Philadelphia, PA, United States (U.S.
 Corporation)
PI US 5470882 19951128
AI US 1994-252716 19940602 (8)
DT Utility
FS Granted
EXNAM Primary Examiner: Dees, Jose G.; Assistant Examiner: Conrad, III, Joseph
 M.
LREP Dinner, Dara L., Venetianer, Stephen, Lentz, Edward T.

CLMN Number of Claims: 5

ECL Exemplary Claim: 1

DRWN No Drawings

LN.CNT 612

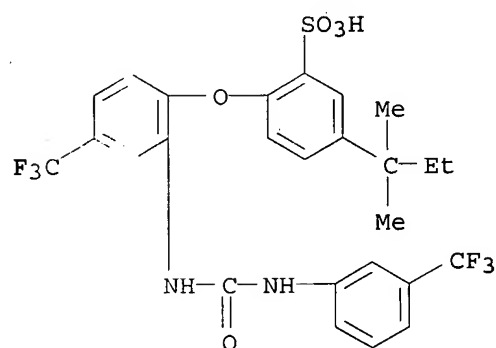
CAS INDEXING IS AVAILABLE FOR THIS PATENT.

IT 447-64-3

(anti-inflammatory benzenesulfonic acid derivs., their prepn., and their activity)

RN 447-64-3 USPATFULL

CN Benzenesulfonic acid, 5-(1,1-dimethylpropyl)-2-[4-(trifluoromethyl)-2-[[[3-(trifluoromethyl)phenyl]amino]carbonyl]amino]phenoxy]- (9CI) (CA INDEX NAME)



L4 ANSWER 12 OF 17 USPATFULL

AB This invention relates to the novel compounds and pharmaceutical compositions of Formula (I).

This invention also relates to a method of treating or reducing inflammation in a mammal in need thereof, which comprises administering to said mammal an effective amount of a compound or composition of Formula (I).

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AN 95:80325 USPATFULL

TI Anti-inflammatory compounds

IN Adams, Jerry L., Wayne, PA, United States

Hall, Ralph F., Villanova, PA, United States

Feibel, George L., Wayne, PA, United States

PA SmithKline Beecham Corp., Philadelphia, PA, United States (U.S. corporation)

PI US 5447957 19950905

AI US 1994-252851 19940602 (8)

DT Utility

FS Granted

EXNAM Primary Examiner: Dees, Jose G.; Assistant Examiner: Barts, Samuel

LREP Ginner, Dara L., Venetianer, Stephen, Lentz, Edward T.

CLMN Number of Claims: 12

ECL Exemplary Claim: 1

DRWN No Drawings

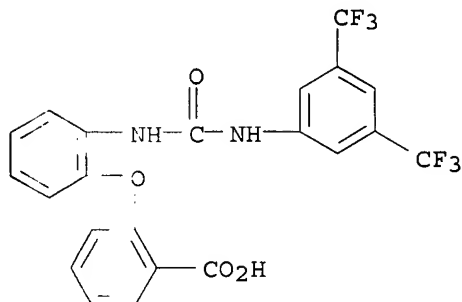
LN.CNT 726

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

IT 171103-10-9P

(antiinflammatory (ureidophenoxy)benzoic acids and derivs. as

inhibitors of phospholipase A2 and CoA-independent transacylase)
 RN 17 103-10-9 USPATFULL
 CN Benzoic acid, 2-[2-[[[3,5-bis(trifluoromethyl)phenyl]amino]carbonyl]amino
 phenoxy]- (9CI) (CA INDEX NAME)

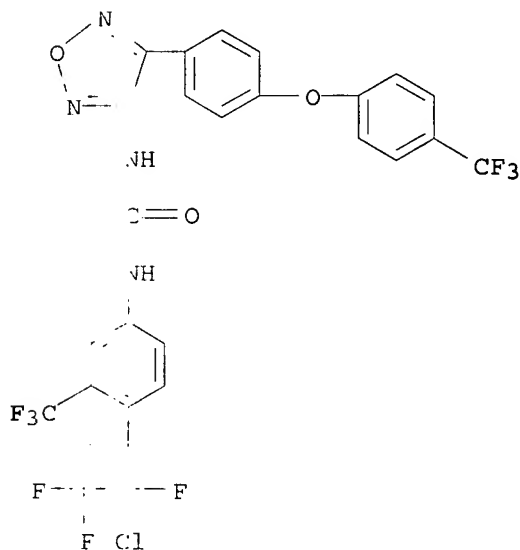


L4 ANSWER 13 OF 17 USPATFULL
 AB Insecticidal and acaricidal novel substituted furazans of the formula
 ##STR1## in which R^{sup.1} and R^{sup.2} are identical or different and
 represent hydrogen, halogen, alkyl, alkoxy, alkylthio, halogenoalkyl,
 halogenoalkoxy, halogenoalkylthio or optionally substituted aryloxy, or
 R^{sup.1} and R^{sup.2} together represent an optionally substituted
 alkylene radical which is interrupted by 1 or 2 oxygen atoms or is
 bonded to the phenyl radical via 1 or 2 oxygen atoms,
 R^{sup.3} and R^{sup.4} are identical or different and represent hydrogen,
 halogen, alkyl, alkoxy, halogenoalkyl or halogenoalkoxy,
 R^{sup.5} represents optionally substituted cycloalkyl, and
 X represents oxygen or sulphur.
 Intermediates of the formula ##STR2## in which B is --NH₂,
 --NO₂ or --NCX,
 R^{sup.3} and R^{sup.4} are identical or different and represent hydrogen,
 halogen, alkyl, alkoxy, halogenoalkyl or halogenoalkoxy,
 R^{sup.6} represents 2,2-difluoro-1-methylcycloprop-1-yl or the ##STR3##
 radical, Y represents hydrogen, methyl, fluorine or chlorine, and
 R^{sup.1} and Y^{sup.2} are identical or different and represent fluorine or
 chlorine,
 are also new.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

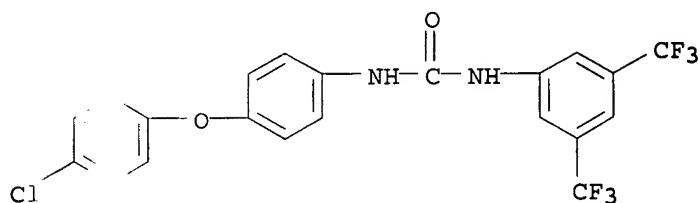
AN 39:63034 USPATFULL
 TI Substituted furazans and insecticidal and acaricidal use
 IN Birrenberg, Wilhelm, Sprockhovel, Germany, Federal Republic of
 Jarhold, Albrecht, Leverkusen, Germany, Federal Republic of
 Steffens, Robert, Cologne, Germany, Federal Republic of
 PA Bayer Aktiengesellschaft, Leverkusen, Germany, Federal Republic of
 (non-U.S. corporation)

PI JS 4853397 19890801
 AI JS 1987-66920 19870625 (7)
 PRAI DE 1986-3622862 19860708
 DT Utility
 FS Granted
 EXNAM Primary Examiner: Raymond, Richard L.
 LREP Sprung Horn Kramer & Woods
 CLMN Number of Claims: 9
 ECL Exemplary Claim: 1,8
 DRWN 10 Drawings
 LN.CNT 174
 CAS INDEXING IS AVAILABLE FOR THIS PATENT.
 IT 113664-71-4P
 (prepn. of, as insecticide and acaricide)
 RN 113664-71-4 USPATFULL
 CN Urea, N-[4-(2-chloro-2,3,3-trifluorocyclobutyl)-3-(trifluoromethyl)phenyl]-
 N'-[4-[4-[4-(trifluoromethyl)phenoxy]phenyl]-1,2,5-oxadiazol-3-yl]-
 (9CI) (CA INDEX NAME)

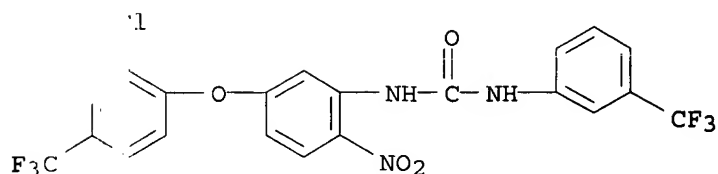


L4 ANSWER 14 OF 17 USPATFULL
 AB The present invention is directed to novel anticoccidial compositions and methods of employing the same to control coccidiosis in poultry. These compositions comprise a polyether antibiotic and a second component which is a selected carbanilide.
 CAS INDEXING IS AVAILABLE FOR THIS PATENT.
 AN 3538961 USPATFULL
 TI Anticoccidial combinations comprising polyether antibiotics and carbanilides
 IN O'Doherty, George O. P., Greenfield, IN, United States
 Clinton, Albert J., Indianapolis, IN, United States
 PI US 4526997 19850702
 AI US 1984-611780 19840518 (6)
 RLI Division of Ser. No. US 1981-260962, filed on 6 May 1981, now patented, Pat. No. US 4468380 which is a continuation of Ser. No. US 1979-107304,

filed on 26 Dec 1979, now abandoned
DT Utility
FS Granted
EXNAM Primary Examiner: Warren, Charles F.; Assistant Examiner: Picard, R. A.
LREP Page, Kathleen R. S., Whale, Arthur R.
CLMN Number of Claims: 12
ECL Exemplary Claim: 1
DRWN No Drawings
LN.CNT 384
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
IT 2063-69-6
(anticoccidal compns. contg. polyether antibiotics and)
RN 2063-69-6 USPATFULL
CN Urea, N-[3,5-bis(trifluoromethyl)phenyl]-N'-[4-(4-chlorophenoxy)phenyl]-
(9CI) (CA INDEX NAME)



L4 ANSWER 15 OF 17 USPATFULL
AB 1,3,5-Triazinones of the formula ##STR1## where R.sup.1, R.sup.2 and R.sup.3 have the meanings given in the description, are used for controlling undesirable plant growth.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
AN 35:23703 USPATFULL
TI 1,3,5-Triazinones and their use for controlling undesirable plant growth
IN Parg, Adolf, Bad Dürkheim, Germany, Federal Republic of
Hamprecht, Gerhard, Weinheim, Germany, Federal Republic of
Wuerzer, Bruno, Otterstadt, Germany, Federal Republic of
PA BASF Aktiengesellschaft, Germany, Federal Republic of (non-U.S. corporation)
PI US 4512797 19850423
AI JS 1983-462024 19830128 (6)
RLI Continuation-in-part of Ser. No. US 1982-446064, filed on 1 Dec 1982, now abandoned
PRAI DE 1981-3147879 19811203
DT Utility
FS Granted
EXNAM Primary Examiner: Ford, John M.
LREP Keil & Weinkauff
CLMN Number of Claims: 8
ECL Exemplary Claim: 1,8
DRWN No Drawings
LN.CNT 300
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
IT 86507-45-6
(cyclocondensation of, with acyl isocyanates)
RN 86507-45-6 USPATFULL
CN Urea, N-[5-[2-chloro-4-(trifluoromethyl)phenoxy]-2-nitrophenyl]-N'-[3-(trifluoromethyl)phenyl]- (9CI) (CA INDEX NAME)



L4 ANSWER 16 OF 17 USPATFULL

AB The present invention is directed to novel anticoccidial compositions and methods of employing the same to control coccidiosis in poultry. These compositions comprise a polyether antibiotic and a second component which is a selected carbanilide.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AI 94:48395 USPATFULL

T Anticoccidial combinations comprising polyether antibiotics and carbanilides

IN O'Doherty, George O. P., Greenfield, IN, United States

Clinton, Albert J., Indianapolis, IN, United States

PA Eli Lilly and Company, Indianapolis, IN, United States (U.S. corporation)

PI JS 4468380 19840828

AI JS 1981-260962 19810506 (6)

RLI Continuation of Ser. No. US 1979-107304, filed on 26 Dec 1979, now abandoned

DT Utility

FS Granted

EXNAM Primary Examiner: Rosen, Sam

LREP Page, Kathleen R. S., Whale, Arthur R.

CLMN Number of Claims: 52

ECL Exemplary Claim: 1,27

DRWN No Drawings

LN.CN. 1366

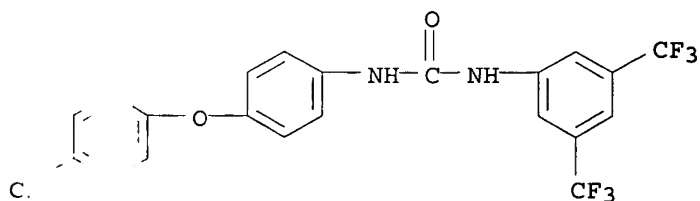
CAS INDEXING IS AVAILABLE FOR THIS PATENT.

IT 2053-69-6

(anticoccidial compns. contg. polyether antibiotics and)

RI 2053-69-6 USPATFULL

CH 2053-69-6 N-[3,5-bis(trifluoromethyl)phenyl]-N'-[4-(4-chlorophenoxy)phenyl]- (9CI) (CA INDEX NAME)



L4 ANSWER 17 OF 17 USPATFULL

AB The present invention is directed to novel anticoccidial compositions and methods of employing the same to control coccidiosis in poultry. These compositions comprise a polyether antibiotic and a second

component selected from nicarbazin and 4,4'-dinitrocarbanilide.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AN 80:40562 USPATFULL
TI Anticoccidial combinations comprising nicarbazin and the polyether
antibiotics
IN Callender, Maurice E., Indianapolis, IN, United States
Jeffers, Thomas K., Greenfield, IN, United States
P Eli Lilly and Company, Indianapolis, IN, United States (U.S.
corporation)
PI US 4218438 19800819
AI US 1979-12165 19790214 (6)
DT Utility
FS Granted
EXNAM Primary Examiner: Rosen, Sam
LREP Page, Kathleen R. S., Whale, Arthur R.
CLMN Number of Claims: 33
ECL Exemplary Claim: 1
DRAW No Drawings
LICNT 352

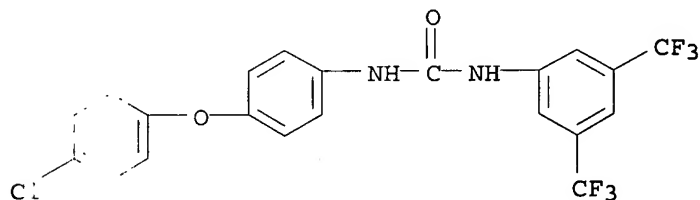
CAS INDEXING IS AVAILABLE FOR THIS PATENT.

IT 2063-69-6

(anticoccidial compn. contg. polyether antibiotic and)

RN 2063-69-6 USPATFULL

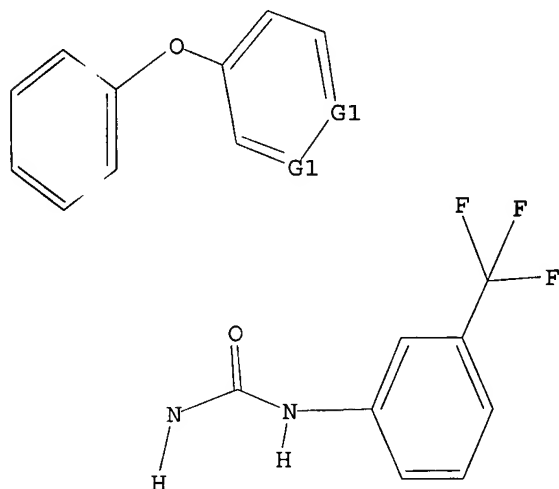
CN 2063-69-6, N-[3,5-bis(trifluoromethyl)phenyl]-N'-[4-(4-chlorophenoxy)phenyl]-
(9CI) (CA INDEX NAME)



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L1 STRUCTURE UPLOADED

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L1 HAS NO ANSWERS
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100.0% PROCESSED 36 ITERATIONS 13 ANSWERS
SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**
BATCH **COMPLETE**
PROJECTED ITERATIONS: 361 TO 1079
PROJECTED ANSWERS: 44 TO 476

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SEARCH TIME: 00.00.02

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=> file uspatall

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FULL ESTIMATED COST	140.28	140.49

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CA INDEXING COPYRIGHT (C) 2002 AMERICAN CHEMICAL SOCIETY (ACS)

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CA INDEXING COPYRIGHT (C) 2002 AMERICAN CHEMICAL SOCIETY (ACS)

=> s 3

L4 17 L3

=> d .bs bib fhitstr 1-17

L4 ANSWER 1 OF 17 USPATFULL

AB This invention relates to the use of a group of heteroaryl ureas containing nitrogen in treating p38 mediated diseases, and pharmaceutical compositions for use in such therapy.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AN 2002:126779 USPATFULL

TI Heteroaryl ureas containing nitrogen hetero-atoms as p38 kinase inhibitors

IN Dumas, Jacques, Orange, CT, UNITED STATES
Riedl, Bernd, Wuppertal, GERMANY, FEDERAL REPUBLIC OF
Khire, Uday, Hamden, CT, UNITED STATES
Sibley, Robert N., North Haven, CT, UNITED STATES
Hatoum-Mokdad, Holia, Hamden, CT, UNITED STATES
Monahan, Mary-Katherine, Hamden, CT, UNITED STATES
Gunn, David E., Hamden, CT, UNITED STATES
Lowinger, Timothy B., Nishinomiya City, JAPAN
Scott, William J., Guilford, CT, UNITED STATES
Smith, Roger A., Madison, CT, UNITED STATES
Wood, Jill E., Hamden, CT, UNITED STATES

PA BAYER CORPORATION (U.S. corporation)

PI US 2002065296 A1 20020530

AI US 2001-838286 A1 20010420 (9)

RLI Continuation-in-part of Ser. No. US 2001-778039, filed on 7 Feb 2001,
PENDING Continuation-in-part of Ser. No. US 1999-425229, filed on 22 Oct
1999, PENDING Continuation of Ser. No. US 1999-257265, filed on 25 Feb
1999, ABANDONED

PRAI US 1999-115878P 19990113 (60)

DT Utility

FS APPLICATION

LREP MILLEN, WHITE, ZELANO & BRANIGAN, P.C., 2200 CLARENDON BLVD., SUITE
1400, ARLINGTON, VA, 22201

CLMN Number of Claims: 39

ECL Exemplary Claim: 1

DRWN No Drawings

LN.CN 2826

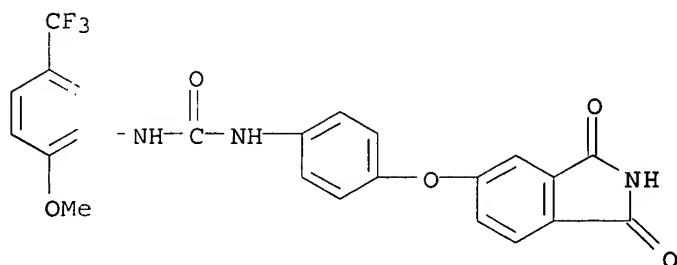
CAS INDEXING IS AVAILABLE FOR THIS PATENT.

IT 284461-54-7P, N-[2-Methoxy-5-(trifluoromethyl)phenyl]-N'-[4-(1,3-dioxoisindolin-5-yl)oxy]phenyl]urea
(prepn. of heteroaryl ureas contg. nitrogen hetero-atoms as p38 kinase inhibitors)

RN 284461-54-7 USPATFULL

CN 'urea, N-[4-[(2,3-dihydro-1,3-dioxo-1H-isoindol-5-yl)oxy]phenyl]-N'-[2-methoxy-5-(trifluoromethyl)phenyl]- (9CI) (CA INDEX NAME)

Print selected from Online session17:01Page 2



L4 ANSWER 2 OF 17 USPATFULL

AB This invention relates to the use of a group of aryl ureas in treating raf mediated diseases, and pharmaceutical compositions for use in such therapy.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AN 2002:78859 USPATFULL

TI Omega-carboxyaryl substituted diphenyl ureas as raf kinase inhibitors

IN Uday, Khire, Hamden, CT, UNITED STATES

Dumas, Jacques, Orange, CT, UNITED STATES

Riedl, Bernd, Wuppertal, GERMANY, FEDERAL REPUBLIC OF

Lowinger, Timothy B., Nishinomiya City, JAPAN

Scott, William J., Guilford, CT, UNITED STATES

Smith, Roger A., Madison, CT, UNITED STATES

Wood, Jill E., Hamden, CT, UNITED STATES

Monahan, Mary-Katherine, Hamden, CT, UNITED STATES

Natero, Reina, Hamden, CT, UNITED STATES

Joel, Renick, Milford, CT, UNITED STATES

Sibley, Robert N., North Haven, CT, UNITED STATES

PA BAYER CORPORATION, Pittsburgh, PA, 15205 (U.S. corporation)

PI US 2002042517 A1 20020411

AI US 2001-948915 A1 20010910 (9)

RLI Continuation of Ser. No. US 1999-425228, filed on 22 Oct 1999, ABANDONED
Continuation-in-part of Ser. No. US 1999-257266, filed on 25 Feb 1999,
ABANDONED

PRAI US 1999-115877P 19990113 (60)

DT Utility

FS APPLICATION

LREP MILLEN, WHITE, ZELANO & BRANIGAN, P.C., 2200 CLARENDON BLVD., SUITE
1400, ARLINGTON, VA, 22201

CLMN Number of Claims: 67

ECL Exemplary Claim: 1

DRWN No Drawings

LN.CN 3675

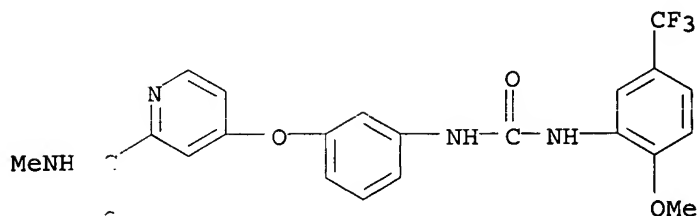
CAS INDEXING IS AVAILABLE FOR THIS PATENT.

IT 2-4461-42-3P

(prepn. of .omega.-carboxy(hetero)aryl substituted di-Ph urea raf
kinase inhibitors by reacting arylisocyanates with arylamines)

RN 2-4461-42-3 USPATFULL

CN 2-Pyridinecarboxamide, 4-[3-[[[2-methoxy-5-(trifluoromethyl)phenyl]amino]
carbonyl]amino]phenoxy]-N-methyl- (9CI) (CA INDEX NAME)



L4 ANSWER 3 OF 17 USPATFULL

AB Chemical structures have been identified which allosterically modify pyruvate kinase and inhibit enzymatic activity. These compounds can be used as pharmaceuticals in the treatment of a wide variety of diseases and disorders where influencing metabolic processes is beneficial, such as the glycolytic pathway, all pathways which use ATP as an energy source, and all pathways which involve 2,3-diphosphoglycerate related to the delivery of oxygen by modifying hemoglobin's oxygen affinity, treatments of tumor and cancer and Alzheimer's disease (AD).

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AN 2001:218507 USPATFULL

TI Allosteric inhibitors of pyruvate kinase

IN Abraham, Donald J., Midlothian, VA, United States

Wang, Changqing, Richmond, CA, United States

Danso-Danquah, Richmond, VA, United States

Burnett, James C., Ashland, VA, United States

Joshi, Gajanan S., Glen Allen, VA, United States

Hoffman, Steven J., Carlisle, MA, United States

PI US 2001046997 A1 20011129

AI US 2001-799873 A1 20010307 (9)

RLI Continuation-in-part of Ser. No. US 1998-46643, filed on 24 Mar 1998, GRANTED, Pat. No. US 6214879

DT Utility

FS APPLICATION

LREP McGuire Woods, LLP, Suite 1800, 1750 Tysons Boulevard, Tysons Corner, McLean, VA, 22102

CLMN Number of Claims: 24

ECL Exemplary Claim: 1

DRWN 7 Drawing Page(s)

LN.CNT 688

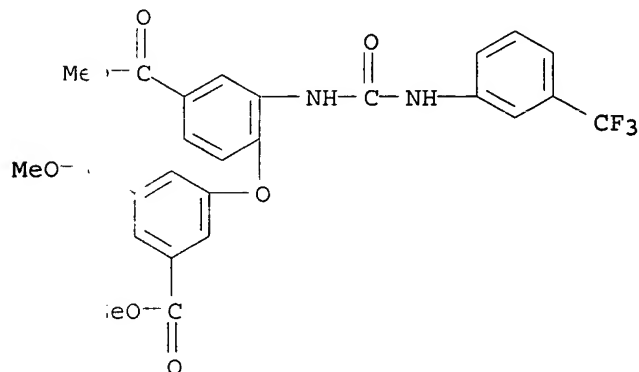
CAS INDEXING IS AVAILABLE FOR THIS PATENT.

IT 289060-07-7

(pyruvate kinase allosteric inhibitors for therapeutic use)

RN 89060-07-7 USPATFULL

CN 3-Benzenedicarboxylic acid, 5-[4-(methoxycarbonyl)-2-[[[3-(trifluoromethyl)phenyl]amino]carbonyl]amino]phenoxy]-, dimethyl ester (9CI) (CA INDEX NAME)



L4 ANSWER 4 OF 17 USPATFULL

AB This invention relates to the use of a group of aryl ureas in treating raf mediated diseases, and pharmaceutical compositions for use in such therapy.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AN 2001:188813 USPATFULL

TI Omega-carboxyaryl substituted diphenyl ureas as raf kinase inhibitors

IN Riedl, Bernd, Wuppertal, Germany, Federal Republic of

Dumas, Jacques, Orange, CT, United States

Khire, Uday, Hamden, CT, United States

Lowinger, Timothy P., Nashomya City, Japan

Scott, William J., Guilford, CT, United States

Smith, Roger A., Madison, CT, United States

Wood, Jill E., Hamden, CT, United States

Monahan, Mary-Katherine, Hamden, CT, United States

Natero, Rena, Hamden, CT, United States

Renick, Joel, Milford, CT, United States

Sibley, Robert N., North Haven, CT, United States

PI US 2001034447 A1 20011025

AI US 2001-773604 A1 20010202 (9)

RLI Continuation of Ser. No. US 1999-425228, filed on 22 Oct 1999, PENDING
Continuation-in-part of Ser. No. US 1999-257266, filed on 25 Feb 1999,
ABANDONED

PRAI US 1999-115877P 19990113 (60)

DT Utility

FS APPLICATION

LREP MILLEN, WHITE, ZELANO & BRANIGAN, P.C., 2200 CLARENDON BLVD., SUITE
1400, ARLINGTON, VA, 22201

CLMN Number of Claims: 67

ECL Exemplary Claim: 1

DRWN No Drawings

LN.CNT 3666

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

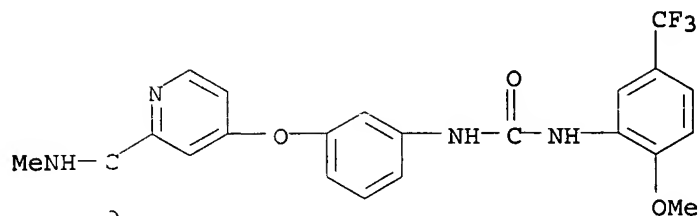
IT 284161-42-3P

(prepn. of .omega.-carboxy(hetero)aryl substituted di-Ph urea raf
kinase inhibitors by reacting arylisocyanates with arylamines)

RN 284161-42-3 USPATFULL

CN 2-Pyridinecarboxamide, 4-[3-[[[2-methoxy-5-(trifluoromethyl)phenyl]amino]
carbonyl]amino]phenoxy]-N-methyl- (9CI) (CA INDEX NAME)

applicants



L4 ANSWER 5 OF 17 USPATFULL

AB This invention relates to the use of a group of aryl ureas in treating raf mediated diseases, and pharmaceutical compositions for use in such therapy.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AN 2001:171152 USPATFULL

TI Omega-carboxyaryl substituted disphenyl ureas as raf kinase inhibitors

IN Riedl, Bernd, Wuppertal, Germany, Federal Republic of

Dumas, Jaques, Orange, CT, United States

Khire, Uday, Hamden, CT, United States

Lowinger, Timothy B., Nishinomiya City, Japan

Scott, William J., Guilford, CT, United States

Smith, Roger A., Madison, CT, United States

Wood, Jill E., Hamden, CT, United States

Monahan, Mary-Katherine, Hamden, CT, United States

Natero, Reina, Hamden, CT, United States

Renick, Joel, Milford, CT, United States

Sibley, Robert N., Noth Haven, CT, United States

PI US 2001027202 A1 20011004

AI US 2001-773658 A1 20010202 (9)

RLI Continuation of Ser. No. US 1999-425228, filed on 22 Oct 1999, PENDING
Continuation-in-part of Ser. No. US 1999-257266, filed on 25 Feb 1999,
ABANDONED

PRAI US 1999-115877P 19990113 (60)

DT Utility

FS APPLICATION

LREP MILLEN, WHITE, ZELANO & BRANIGAN, P.C., Arlington Courthouse Plaza I,
Suite 1400, 2200 Clarendon Boulevard, Arlington, VA, 22201

CLMN Number of Claims: 67

ECL Exemplary Claim: 1

DRWN No Drawings

LN.CN 3656

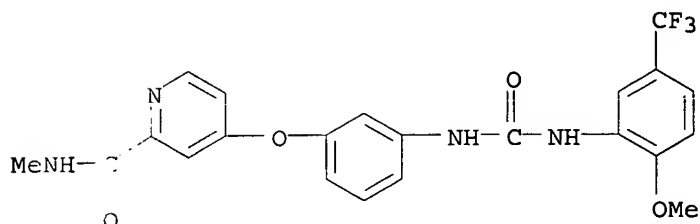
CAS INDEXING IS AVAILABLE FOR THIS PATENT.

IT 284461-42-3P

(prepn. of .omega.-carboxy(hetero)aryl substituted di-Ph urea raf
kinase inhibitors by reacting arylisocyanates with arylamines)

RN 284461-42-3 USPATFULL

CN 2-Pyridinecarboxamide, 4-[3-[[[2-methoxy-5-(trifluoromethyl)phenyl]amino]
carbonyl]amino]phenoxy]-N-methyl- (9CI) (CA INDEX NAME)



L4 ANSWER 6 OF 17 USPATFULL

AB This invention relates to the use of a group of aryl ureas in treating raf mediated diseases, and pharmaceutical compositions for use in such therapy.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AN 2001:139616 USPATFULL

TI Omega-carboxyaryl substituted diphenyl ureas as raf kinase inhibitors

IN Fiedl, Bernd, Wupperal, Germany, Federal Republic of

Dumas, Jacques, Orange, CT, United States

Khire, Uday, Hamden, CT, United States

Lowinger, Timothy B., Nashnomya City, Japan

Scott, William J., Guilford, CT, United States

Smith, Roger A., Madison, CT, United States

Wood, Jill E., Hamden, CT, United States

Monahan, Mary-Katherine, Hamden, CT, United States

Natero, Rena, Hamden, CT, United States

Renick, Joel, Milford, CT, United States

Sibley, Robert N., North Haven, CT, United States

PI US 2001016659 A1 20010823

AI US 2001-773672 A1 20010202 (9)

RLI Continuation of Ser. No. US 1999-425228, filed on 22 Oct 1999, PENDING
Continuation-in-part of Ser. No. US 1999-257266, filed on 25 Feb 1999,
ABANDONED

PRAI US 1999-115877P 19990113 (60)

DT Utility

FS APPLICATION

LREP MILLEN, WHITE, ZELANO & BRANIGAN, P.C., 2200 CLARENDON BLVD., SUITE
1400, ARLINGTON, VA, 22201

CLMN Number of Claims: 67

ECL Exemplary Claim: 1

DRWN No Drawings

LN.CNT 3652

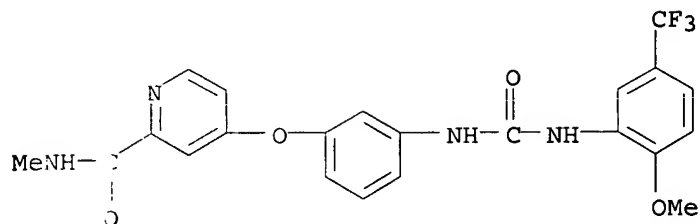
CAS INDEXING IS AVAILABLE FOR THIS PATENT.

IT 284461-42-3P

(prepn. of omega.-carboxy(hetero)aryl substituted di-Ph urea raf
kinase inhibitors by reacting arylisocyanates with arylamines)

RN 284461-42-3 USPATFULL

CN 2-Pyridinecarboxamide, 4-[3-[[[2-methoxy-5-(trifluoromethyl)phenyl]amino]
carbonyl]amino]phenoxy]-N-methyl- (9CI) (CA INDEX NAME)



L4 ANSWER 7 OF 17 USPATFULL

AB This invention relates to the use of a group of aryl ureas in treating raf mediated diseases, and pharmaceutical compositions for use in such therapy.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AN 2001:123628 USPATFULL

TI omega-carboxyaryl substituted diphenyl ureas as raf kinase inhibitors

IN Riedl, Bernd, Wuppertal, Germany, Federal Republic of

Dumas, Jacques, Orange, CT, United States

Khire, Uday, Hamden, CT, United States

Lowinger, Timothy B., Nishinomiya City, Japan

Scott, William J., Guilford, CT, United States

Smith, Roger A., Madison, CT, United States

Wood, Jill E., Hamden, CT, United States

Monahan, Mary-Katherine, Hamden, CT, United States

Natero, Reina, Hamden, CT, United States

Renick, Joel, Milford, CT, United States

Sibley, Robert N., North Haven, CT, United States

PI US 2001011136 A1 20010802

AI US 2001-773675 A1 20010202 (9)

RLI Continuation of Ser. No. US 1999-425228, filed on 22 Oct 1999, PENDING
Continuation-in-part of Ser. No. US 1999-257266, filed on 25 Feb 1999,
ABANDONED

PRAI US 1999-115877P 19990113 (60)

DT Utility

FS APPLICATION

LREP MILLEN, WHITE, ZELANO & BRANIGAN, P.C., Suite 1400, 2200 Clarendon
Blvd., Arlington, VA, 22201

CLMN Number of Claims: 67

ECL Exemplary Claim: 1

DRWN No Drawings

LN.CN 3646

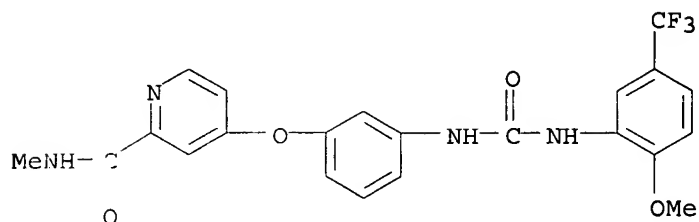
CAS INDEXING IS AVAILABLE FOR THIS PATENT.

IT 284461-42-3P

(prepn. of omega-carboxy(hetero)aryl substituted di-Ph urea raf
kinase inhibitors by reacting arylisocyanates with arylamines)

RN 284461-42-3 USPATFULL

CN 2-Pyridinecarboxamide, 4-[3-[[[2-methoxy-5-(trifluoromethyl)phenyl]amino]
carbonyl]amino]phenoxy]-N-methyl- (9CI) (CA INDEX NAME)



L4 ANSWER 8 OF 17 USPATFULL

AB This invention relates to the use of a group of aryl ureas in treating raf mediated diseases, and pharmaceutical compositions for use in such therapy.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AN 2001:123627 USPATFULL

TI Omega-carboxyaryl substituted diphenyl ureas as raf kinase inhibitors

IN Riedl, Bernd, Wuppertal, Germany, Federal Republic of

Dumas, Jacques, Orange, CT, United States

Khire, Uday, Hamden, CT, United States

Lowinger, Timothy B., Nishinomiya City, Japan

Scott, William J., Guilford, CT, United States

Smith, Roger A., Madison, CT, United States

Wood, Jill E., Hamden, CT, United States

Monahan, Mary-Katherine, Hamden, CT, United States

Natero, Reina, Hamden, CT, United States

Renick, Joel, Milford, CT, United States

Sibley, Robert N., North Haven, CT, United States

PI US 2001011135 A1 20010802

AI US 2001-773659 A1 20010202 (9)

RLI Continuation of Ser. No. US 1999-425228, filed on 22 Oct 1999, PENDING
Continuation-in-part of Ser. No. US 1999-257266, filed on 25 Feb 1999,
ABANDONED

PRAI US 1999-115877P 19990113 (60)

DT Utility

FS APPLICATION

LREP MILLEN, WHITE, ZELANO & BRANIGAN, P.C., Suite 1400, Arlington Courthouse
Plaza 1, Arlington, VA, 22201

CLMN Number of Claims: 67

ECL Exemplary Claim: 1

DRWN No Drawings

LN.CN 3686

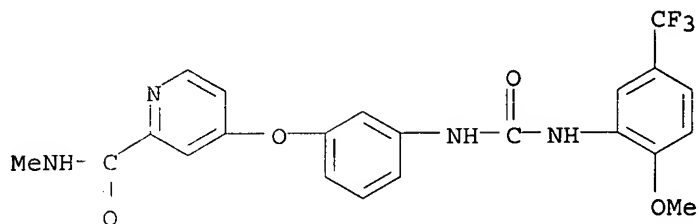
CAS INDEXING IS AVAILABLE FOR THIS PATENT.

IT 284461-42-3P

(prepn. of .omega.-carboxy(hetero)aryl substituted di-Ph urea raf
kinase inhibitors by reacting arylisocyanates with arylamines)

RN 284461-42-3 USPATFULL

CN 2-Pyridinecarboxamide, 4-[3-[[[2-methoxy-5-(trifluoromethyl)phenyl]amino]
carbonyl]amino]phenoxy]-N-methyl- (9CI) (CA INDEX NAME)



L4 ANSWER 9 OF 17 USPATFULL

AB The present invention relates to novel quinoline derivatives and quinazoline derivatives represented by the following formula (I):
 ##STR1## [wherein R.sub.1 and R.sub.2 are each independently H or C.sub.1 -C.sub.4 -alkyl, or R.sub.1 and R.sub.2 together form C.sub.1 -C.sub.3 -alkylene, X is O, S or CH.sub.2, W is CH or N, and Q is a substituted aryl group or substituted heteroaryl group] and their pharmaceutically acceptable salts, having platelet-derived growth factor receptor autophosphorylation inhibitory activity, to pharmaceutical compositions containing these compounds, and to methods for the treatment of diseases associated with abnormal cell growth such as tumors.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AN 2000:150184 USPATFULL

TI Quinoline and quinazoline derivatives inhibiting platelet-derived growth factor receptor autophosphorylation and pharmaceutical compositions containing the same

IN Kubo, Kazuo, Takasaki, Japan
 Ohyama, Shinichi, Takasaki, Japan
 Shimizu, Toshiyuki, Takasaki, Japan
 Nishitoba, Tsuyoshi, Takasaki, Japan
 Kato, Shinichiro, Takasaki, Japan
 Murooka, Hideko, Takasaki, Japan
 Kobayashi, Yoshiko, Takasaki, Japan

PA Kirin Beer Kabushiki Kaisha, Tokyo-to, Japan (non-U.S. corporation)

PI US 6143764 20001107

WO 9717329 19970515

AI US 1998-68660 19980506 (9)

WO 1996-JP3229 19961105

19980506 PCT 371 date

19980506 PCT 102(e) date

PRAI JP 1995-313555 19951107

JP 1996-62121 19960223

DT Utility

FS Granted

EXNAM Primary Examiner: Seaman, D. Margaret

LREP Foley & Lardner

CLMN Number of Claims: 52

ECL Exemplary Claim: 1

DRWN No Drawings

LN.CNT 5569

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

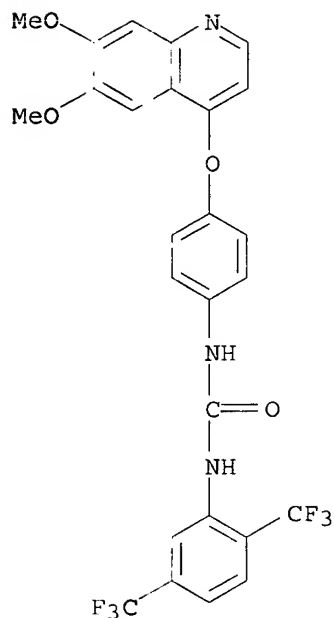
IT 190727-78-7P

(prepn. of quinoline and quinazoline derivs. inhibiting platelet-derived growth factor receptor autophosphorylation)

RN 190727-78-7 USPATFULL

CN Urea, N-[2,5-bis(trifluoromethyl)phenyl]-N'-[4-[(6,7-dimethoxy-4-

quinolinyl]oxy]phenyl] - (9CI) (CA INDEX NAME)



L4 ANSWER 10 OF 17 USPATFULL

AB This invention relates to the novel pharmaceutical compositions of Formulas (I) and (II) each of which comprises a compound of Formula (I) or (II) and a pharmaceutically acceptable diluent or carrier. This invention also relates to a method of treating or reducing inflammation in a mammal in need thereof, which comprises administering to said mammal an effective amount of a compound or composition of Formula (I) or (II).

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AN 1999:67289 USPATFULL

TI Anti-inflammatory compounds

IN Dixon, James Scott, Malvern, PA, United States

Hall, Ralph Floyd, Villanova, PA, United States

Marshall, Lisa Ann, Wyndmoor, PA, United States

Chilton, III, Floyd H., Pilot Mountain, NC, United States

Mayer, Ruth Judik, Wayne, PA, United States

Winkler, James David, Fort Washington, PA, United States

PA SmithKline Beecham Corporation, Philadelphia, PA, United States (U.S. corporation)

The Johns Hopkins University, Baltimore, MD, United States (U.S. corporation)

PI US 5912270 19990615

WO 9533712 19951214

AI US 1996-737650 19961122 (8)

WO 1995-US6677 19950602

19961122 PCT 371 date

19961122 PCT 102(e) date

RLI Continuation-in-part of Ser. No. US 1994-252716, filed on 2 Jun 1994, now patented, Pat. No. US 5470882

DT Utility

FS Granted

EXNAM Primary Examiner: Gerstl, Robert

LREP Dinner, Dara L., Venetianer, Stephen, Kinzig, Charles

CLMN Number of Claims: 15

ECL Exemplary Claim: 1

DRWN No Drawings

LN.CNT 1767

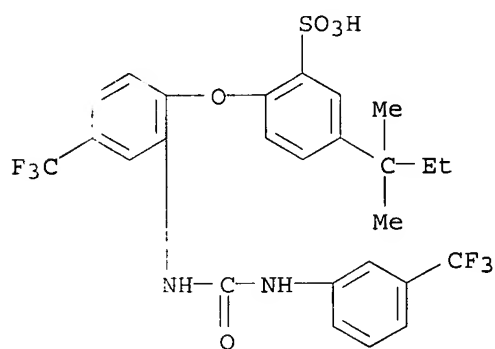
CAS INDEXING IS AVAILABLE FOR THIS PATENT.

IT 447-64-3P

(prepn. of antiinflammatory ureidophenoxybenzenesulfonates)

RN 447-64-3 USPATFULL

CN Benzenesulfonic acid, 5-(1,1-dimethylpropyl)-2-[4-(trifluoromethyl)-2-[[[3-(trifluoromethyl)phenyl]amino]carbonyl]amino]phenoxy] - (9CI) (CA INDEX NAME)



L4 ANSWER 11 OF 17 USPATFULL

AB This invention relates to the novel pharmaceutical compositions of Formulas (I) and (II) each of which comprises a compound of Formula (I) or (II) and a pharmaceutically acceptable diluent or carrier.

This invention also relates to a method of treating or reducing inflammation in a mammal in need thereof, which comprises administering to said mammal an effective amount of a compound or composition of Formula (I) or (II).

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AN 95:105872 USPATFULL

TI Anti-inflammatory compounds

IN Dixon, James S., Malvern, PA, United States

Hall, Ralph F., Villanova, PA, United States

Marshall, Lisa A., Wyndmoor, PA, United States

Chilton, III, Floyd H., Pilot Mountain, NC, United States

Mayer, Ruth J., Wayne, PA, United States

Winkler, James D., Fort Washington, PA, United States

PA SmithKline Beecham Corp., Philadelphia, PA, United States (U.S. corporation)

PI US 5470882 19951128

AI US 1994-252716 19940602 (8)

DT Utility

FS Granted

EXNAM Primary Examiner: Dees, Jose G.; Assistant Examiner: Conrad, III, Joseph M.

LREP Dinner, Dara L., Venetianer, Stephen, Lentz, Edward T.

CMN Number of Claims: 5

ECL Exemplary Claim: 1

DRWN No Drawings

LN.CNT 1612

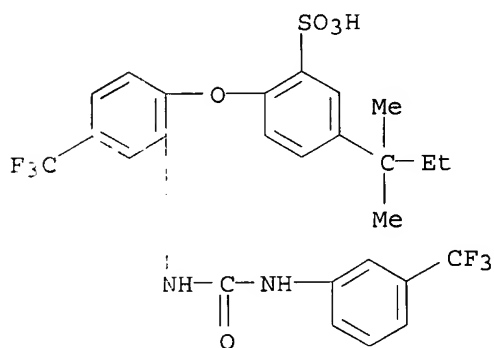
CAS INDEXING IS AVAILABLE FOR THIS PATENT.

IT 447-64-3

(anti-inflammatory benzenesulfonic acid derivs., their prepn., and their activity)

RN 447-64-3 USPATFULL

CN Benzenesulfonic acid, 5-(1,1-dimethylpropyl)-2-[4-(trifluoromethyl)-2-[[[3-(trifluoromethyl)phenyl]amino]carbonyl]amino]phenoxy]- (9CI) (CA INDEX NAME)



L4 ANSWER 12 OF 17 USPATFULL

AB This invention relates to the novel compounds and pharmaceutical compositions of Formula (I).

This invention also relates to a method of treating or reducing inflammation in a mammal in need thereof, which comprises administering to said mammal an effective amount of a compound or composition of Formula (I).

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AN 95:80325 USPATFULL

TI Anti-inflammatory compounds

IN Adams, Jerry L., Wayne, PA, United States

Hall, Ralph F., Villanova, PA, United States

Seibel, George L., Wayne, PA, United States

PA SmithKline Beecham Corp., Philadelphia, PA, United States (U.S. corporation)

PI US 5447957 19950905

AI US 1994-252851 19940602 (8)

DT Utility

FS Granted

EXNAM Primary Examiner: Dees, Jose G.; Assistant Examiner: Barts, Samuel

L EP Danner, Dara L., Venetianer, Stephen, Lentz, Edward T.

CMN Number of Claims: 12

ECL Exemplary Claim: 1

DPWN No Drawings

LN.CNT 1616

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

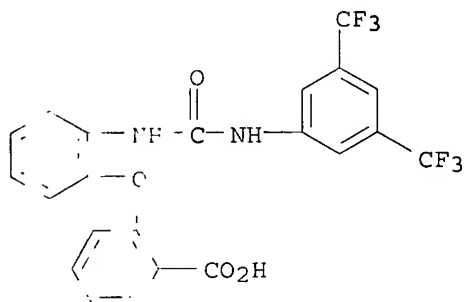
IT 171103-10-9P

(antiinflammatory (ureidophenoxy)benzoic acids and derivs. as

Inhibitors of phospholipase A2 and CoA-independent transacylase)

RI 171103-10-9 USPATFULL

CI Benzoic acid, 2-[2-[[[3,5-bis(trifluoromethyl)phenyl]amino]carbonyl]amino
[phenoxy]- (9CI) (CA INDEX NAME)



LI ANSWER 13 OF 17 USPATFULL

A1 Insecticidal and acaricidal novel substituted furazans of the formula ##STR1## in which R^{sup.1} and R^{sup.2} are identical or different and represent hydrogen, halogen, alkyl, alkoxy, alkylthio, halogenoalkyl, halogenoalkoxy, halogenoalkylthio or optionally substituted aryloxy, or

R^{sup.1} and R^{sup.2} together represent an optionally substituted alkylene radical which is interrupted by 1 or 2 oxygen atoms or is bonded to the phenyl radical via 1 or 2 oxygen atoms,

R^{sup.3} and R^{sup.4} are identical or different and represent hydrogen, halogen, alkyl, alkoxy, halogenoalkyl or halogenoalkoxy,

R^{sup.5} represents optionally substituted cycloalkyl, and

X represents oxygen or sulphur.

Intermediates of the formula ##STR2## in which B is --NH_{sub.2}, --NO_{sub.2} or --NCX,

R^{sup.3} and R^{sup.4} are identical or different and represent hydrogen, halogen, alkyl, alkoxy, halogenoalkyl or halogenoalkoxy,

R^{sup.6} represents 2,2-difluoro-1-methylcycloprop-1-yl or the ##STR3## radical, Y represents hydrogen, methyl, fluorine or chlorine, and

Y^{sup.1} and Y^{sup.2} are identical or different and represent fluorine or chlorine,

are also new.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AI 89:63034 USPATFULL

TJ Substituted furazans and insecticidal and acaricidal use

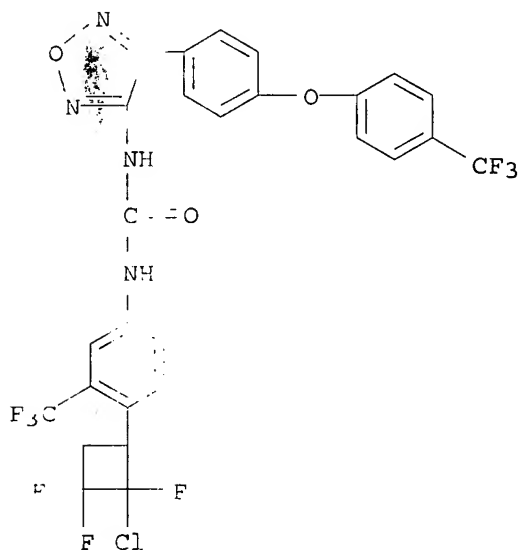
IN Sorenberg, Wilhelm, Sprockhovel, Germany, Federal Republic of

Machold, Albrecht, Leverkusen, Germany, Federal Republic of

Steffens, Robert, Cologne, Germany, Federal Republic of

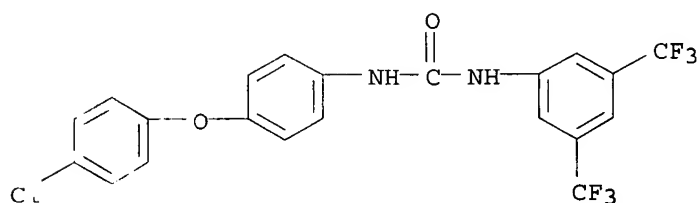
PI Bayer Aktiengesellschaft, Leverkusen, Germany, Federal Republic of (non-U.S. corporation)

PI US 4853397 19890801
 A US 1987-66920 19870625 (7)
 PPAI DE 1986-3622862 19860708
 DT Utility
 FS Granted
 EXNAM Primary Examiner: Raymond, Richard L.
 LREP Sprung Horn Kramer & Woods
 CLMN Number of Claims: 9
 ECL Exemplary Claim: 1,8
 DWN No Drawings
 L1.CNT 874
 CAS INDEXING IS AVAILABLE FOR THIS PATENT.
 I 113664-71-4P
 'prepn. of, as insecticide and acaricide)
 RI 113664-71-4 USPATFULL
 CH Urea, N-[4-(2-chloro-2,3,3-trifluorocyclobutyl)-3-(trifluoromethyl)phenyl]-
 N'-[4-[4-[4-(trifluoromethyl)phenoxy]phenyl]-1,2,5-oxadiazol-3-yl]-
 (3CI) (CA INDEX NAME)

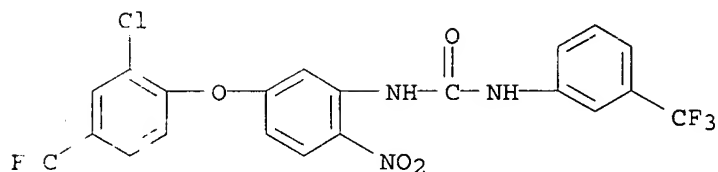


L ANSWER 14 OF 17 USPATFULL
 AB The present invention is directed to novel anticoccidial compositions and methods of employing the same to control coccidiosis in poultry. These compositions comprise a polyether antibiotic and a second component which is a selected carbanilide.
 CAS INDEXING IS AVAILABLE FOR THIS PATENT.
 AN 85:38961 USPATFULL
 T Anticoccidial combinations comprising polyether antibiotics and carbanilides
 I O'Doherty, George O. P., Greenfield, IN, United States
 Cinton, Albert J., Indianapolis, IN, United States
 P US 4526997 19850702
 A US 1984-611780 19840518 (6)
 RI Division of Ser. No. US 1981-260962, filed on 6 May 1981, now patented, Pat. No. US 4468380 which is a continuation of Ser. No. US 1979-107304,

filed on 26 Dec 1979, now abandoned
DT Utility
FS Granted
EXNAM Primary Examiner: Warren, Charles F.; Assistant Examiner: Picard, R. A.
LREP Page, Kathleen R. S., Whale, Arthur R.
CLMN Number of Claims: 12
ECL Exemplary Claim: 1
DWN No Drawings
LFCNT 884
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
IN 2063-69-6
(anticoccidial compns. contg. polyether antibiotics and)
PI 2063-69-6 USPATFULL
CI Urea, N-[3,5-bis(trifluoromethyl)phenyl]-N'-[4-(4-chlorophenoxy)phenyl]-
(9CI) (CA INDEX NAME)



LA ANSWER 15 OF 17 USPATFULL
A 1,3,5-Triazinones of the formula ##STR1## where R.sup.1, R.sup.2 and R.sup.3 have the meanings given in the description, are used for controlling undesirable plant growth.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
AI 85:23703 USPATFULL
TI 1,3,5-Triazinones and their use for controlling undesirable plant growth
IN Parg, Adolf, Bad Dürkheim, Germany, Federal Republic of
Hamprecht, Gerhard, Weinheim, Germany, Federal Republic of
Wuerzer, Bruno, Otterstadt, Germany, Federal Republic of
P BASF Aktiengesellschaft, Germany, Federal Republic of (non-U.S. corporation)
P US 4512797 19850423
AI US 1983-462024 19830128 (6)
RMI Continuation-in-part of Ser. No. US 1982-446064, filed on 1 Dec 1982, now abandoned
PAI DE 1981-3147879 19811203
DT Utility
FS Granted
EXNAM Primary Examiner: Ford, John M.
LREP Keil & Weinkauff
CMN Number of Claims: 8
ECL Exemplary Claim: 1,8
DWN No Drawings
LFCNT 300
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
IN 86607-45-6
(cyclocondensation of, with acyl isocyanates)
RI 86607-45-6 USPATFULL
CI Urea, N-[5-[2-chloro-4-(trifluoromethyl)phenoxy]-2-nitrophenyl]-N'-[3-(trifluoromethyl)phenyl]- (9CI) (CA INDEX NAME)



L ANSWER 16 OF 17 USPATFULL

AB The present invention is directed to novel anticoccidial compositions and methods of employing the same to control coccidiosis in poultry. These compositions comprise a polyether antibiotic and a second component which is a selected carbanilide.

C S INDEXING IS AVAILABLE FOR THIS PATENT.

A 3448395 USPATFULL

T Anticoccidial combinations comprising polyether antibiotics and carbanilides

IN O'Doherty, George O. P., Greenfield, IN, United States

Clinton, Albert J., Indianapolis, IN, United States

P Eli Lilly and Company, Indianapolis, IN, United States (U.S. Corporation)

P US 4468380 19840828

A US 1981-260962 19810506 (6)

P-I Continuation of Ser. No. US 1979-107304, filed on 26 Dec 1979, now abandoned

D Utility

F3 Granted

E'NAM Primary Examiner: Rosen, Sam

L EP Page, Kathleen R. S., Whale, Arthur R.

C MN Number of Claims: 52

E L Exemplary Claim: 1,27

D-WN 10 Drawings

L .CNT 1366

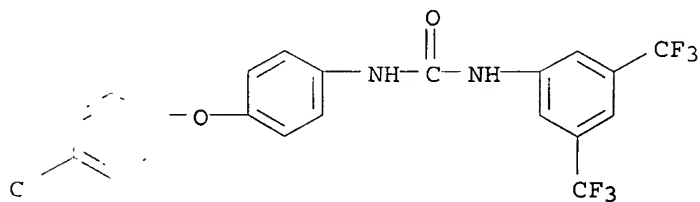
C S INDEXING IS AVAILABLE FOR THIS PATENT.

IN 2063-69-6

Anticoccidial compns. contg. polyether antibiotics and

PY 2063-69-6 USPATFULL

C Urea, N-[3,5-bis(trifluoromethyl)phenyl]-N'-[4-(4-chlorophenoxy)phenyl]- (OCT) (CA INDEX NAME)



L ANSWER 17 OF 17 USPATFULL

A The present invention is directed to novel anticoccidial compositions and methods of employing the same to control coccidiosis in poultry. These compositions comprise a polyether antibiotic and a second

Component selected from nicarbazin and 4,4'-dinitrocarbanilide.

C S INDEXING IS AVAILABLE FOR THIS PATENT.

A : 40562 USPATFULL

T : Anticoccidial combinations comprising nicarbazin and the polyether antibiotics

I : Callender, Maurice E., Indianapolis, IN, United States

Offers, Thomas K., Greenfield, IN, United States

P : Eli Lilly and Company, Indianapolis, IN, United States (U.S.

Corporation)

P : 4218438 19800819

A : 1979-12165 19790214 (6)

D : Utility

F : Granted

E NAM Primary Examiner: Rosen, Sam

L EP Age, Kathleen R. S., Whale, Arthur R.

C MN Number of Claims: 33

E L Exemplary Claim: 1

D WN No Drawings

I .CNT 2

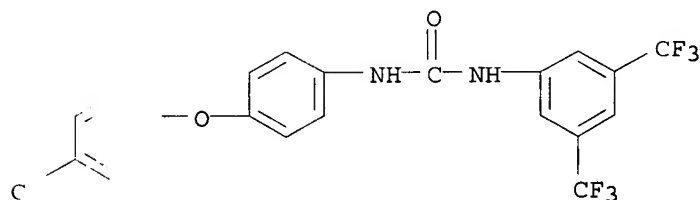
C S INDEXING IS AVAILABLE FOR THIS PATENT.

I 2067-69-6

(anticoccidial compn. contg. polyether antibiotic and)

P 2067-69-6 USPATFULL

C Uracil, N-[3,5-bis(trifluoromethyl)phenyl]-N'-[4-(4-chlorophenoxy)phenyl]-
(CAI) (CA INDEX NAME)



=~ file caplus

C ST IN U.S. DOLLARS

SINCE FILE

TOTAL

ENTRY

SESSION

F LL ESTIMATED COST

102.91

243.40

F LE 'CAPLUS' ENTERED AT 16:58:08 ON 15 JUL 2002

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FILE COVERS 1907 - 15 Jul 2002 VOL 137 ISS 3
FILE LAST UPDATED: 14 Jul 2002 (20020714/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

CAS roles have been modified effective December 16, 2001. Please check your SDI profiles to see if they need to be revised. For information on CAS roles, enter HELP ROLES at an arrow prompt or use the CAS Roles thesaurus (/RL field) in this file.

= d h s

(FILE 'HOME' ENTERED AT 16:55:35 ON 15 JUL 2002)

FILE 'REGISTRY' ENTERED AT 16:55:43 ON 15 JUL 2002

I: STRUCTURE UPLOADED

I: 13 S L1

I: 365 S L1 FUL

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I: 17 S L3

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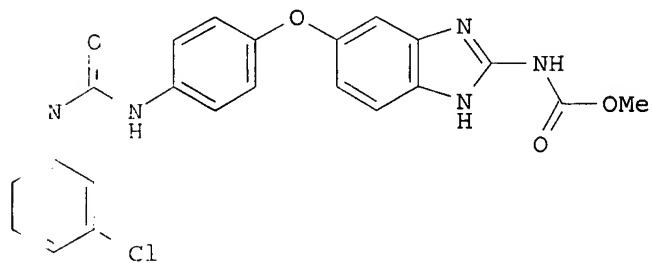
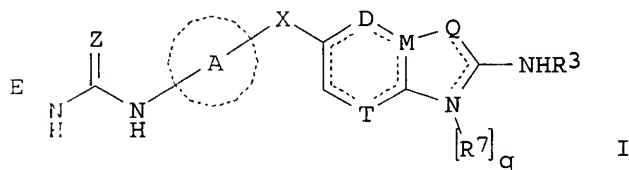
= s l3

I: 32 L3

= d abs bib fhitr 1-32

L: ANSWER 1 OF 32 CAPLUS COPYRIGHT 2002 ACS

G



A: The title compds. [I; E = (un)substituted aryl, heteroaryl; A = aryl,

heteroaryl, heterocyclyl; X = S, O, SO₂, SO, CH₂, CHOH, CO; Z = O, S; p = 0-1; q = 0-1; D = CH, T = CR₈, M = C and Q = NT₇p, wherein p = 0 and q = 1; or D = CH, T = CR₈, M = C and Q = NR₇p, wherein p = 1 and q = 0, or D = CH, T = CR₈, M = C and Q = S or O, wherein q = 0; or D = N, T = CR₈, M = C and Q = NR₇p, wherein either p or q = 0 and the other = 1; or D = CH, T = N, M = C and Q = NR₇p, wherein either p or q = 0 and the other = 1; or D = CH, T = CR₈, M = N and Q = CH, wherein q = 0; R₁ = alkyl, haloalkyl, aryl, etc.; R₂ = H, alkyl, aryl, etc.; R₃ = alkylene or alkylene substituted by oxo, and is linked together with N atom to which it is attached and to one of the benzimidazole N atoms to form a heterocyclic compd. fused to the benzimidazole; R₇ = H, alkyl, etc.; R₈ = H, halo and their salts, useful in the treatment of hyperproliferative diseases, were prepd. Thus, reacting Me [5-(4-aminophenoxy)-1H-benzimidazol-2-yl]carbamate (prepn. given) with 3-chlorophenyl isocyanate in THF afforded 69% II which showed pIC₅₀ of > 7.0 in TIE-2 and VEGFR2 enzyme assays.

A 2002:428885 CAPLUS

D 137:6179

T Preparation of benzimidazoles as TIE-2 and/or VEGFR2 inhibitors

I Cheung, Mui; Harris, Philip Anthony; Hasegawa, Masaichi; Ida, Satoru; Kano, Kazuya; Nishigaki, Naohiko; Sato, Hideyuki; Veal, James Martin; Washio, Yoshiaki; West, Rob I.

PA Glaxo Group Limited, UK; Glaxosmithkline K.K.

SP PCT Int. Appl., 217 pp.

CODEN: PIXXD2

D Patent

L English

F N.CNT 1

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2002044156	A2	20020606	WO 2001-US44553	20011128
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG				

P MI US 2000-253868P P 20001129

US 2001-310939P P 20010808

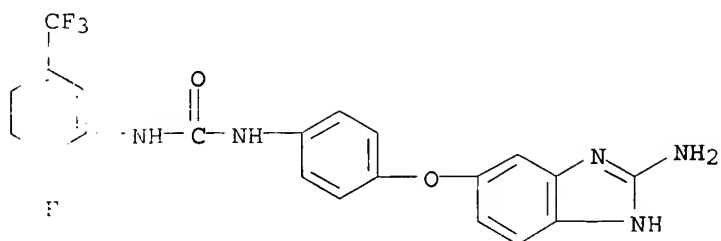
OS MARPAT 137:6179

I 433224-24-9P

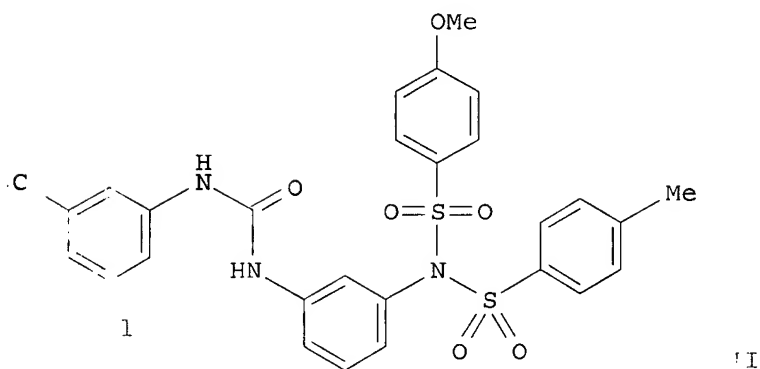
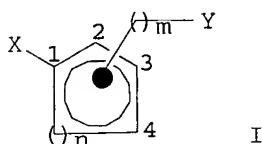
ES: PAC (Pharmacological activity); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses) (prepn. of benzimidazoles as TIE-2 and/or VEGFR2 inhibitors)

P 433224-24-9 CAPLUS

C Mea, N-[4-[(2-amino-1H-benzimidazol-5-yl)oxy]phenyl]-N'-[2-fluoro-5-(trifluoromethyl)phenyl]- (9CI) (CA INDEX NAME)



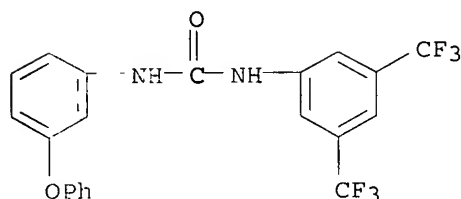
L ANSWER 2 OF 32 CAPLUS COPYRIGHT 2002 ACS
G



A Title compds. I [n = 1-2 forming a central 5-6 membered (un)satd. carbocyclic ring; m = 0-3; [CH₂]_mY is attached to said central carbocyclic ring at position 2, 3, or 4; X, Y = carboxamide, thiocarboxamide, ureido, aminosulfonyl, etc.] were prepd. Examples include over 30 compds. synthesized, assays for rotamase inhibition, neuronal cell growth/regeneration, in-vivo protective effects in an animal model of stroke/myocardial infarction (rat) and an in-vivo model of hair growth (mouse). For instance, 3-nitroaniline was reacted with 4-methylphenylsulfonylsulfonyl chloride and 4-methoxyphenylsulfonyl chloride (DMA, Et₃N) to give the bis(sulfonamide) as a solid. This intermediate was reduced (EtOH/Na, NH₄Cl, In.degree., reflux, 4 h) and subsequently treated with 3,5-dichlorophenylisocyanate to give II. II had IC₅₀ = 162 nM for rotamase (a measure of cyclophilin (CyP) A binding). I have an affinity for CyP-type immunophilin proteins and are useful for the

treatment of neurol. disorders, hair loss disorders, ischemic disorders,
and disorders caused by viral or protozoan infection.
AN 2002:428855 CAPLUS
DN 137:20228
TI Sulfonamido/amido/ureido-phenyl-amides as cyclophilin binding compounds
IN Hamilton, Gregory S.; Belyakov, Sergei; Vaal, Mark; Wei, Ling; Wu,
Yong-Qian; Steiner, Joseph P.
PA Guilford Pharmaceuticals Inc., USA
SO PCT Int. Appl., 141 pp.
CODEN: PIXXD2
DT Patent
LA English
FAN. C 1

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2002044126	A2	20020606	WO 2001-US44449	20011128
<p>W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG</p>				
PRAI US 2000-253074P	P	20001128		
US 2001-291966P	P	20010521		
OS MARPAT 137:20228				
IT 1995-43-3P				
<p>Int. PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (uses) (drug; prepn. of 1,3-disubstituted sulfonamido/amido/ureido-Ph-amides as immunophilin ligands)</p>				
RN 1995-43-3 CAPLUS				
CN Urea, N-[3,5-bis(trifluoromethyl)phenyl]-N'-(3-phenoxyphenyl)- (9CI) (CA INDEX NAME)				



L5 ANSWER 3 OF 32 CAPLUS COPYRIGHT 2002 ACS
AB This invention relates to the use of a group of heteroaryl ureas (I; for
example, N-(2-methoxy-3-quinolyl)-N'-(4-[3-(N-
methylcarbamoyl)phenoxy]phenyl]urea) contg. N in treating p38 mediated
diseases, and pharmaceutical compns. for use in such therapy. I is
A NHC(O)NH-B or a pharmaceutically acceptable salt thereof, wherein A is a
substituted or unsubstituted pyridyl, quinolinyl or isoquinolinyl group, B
is a substituted or unsubstituted, up to tricyclic aryl or heteroaryl
moiety of up to 50 C atoms with a cyclic structure bound directly to N,
contg. at least 5 cyclic members with 0-3 members of groups consisting of

N, O and S. Information about the substituents for A and B are given in the claims. Although the methods of prepn. are not claimed, 37 example prepn. are included as well as examples of prepn. of intermediates. No pharmacol. data is included.

AN 2 02:409267 CAPLUS

DN 137:6098

TI Heteroaryl ureas containing nitrogen hetero-atoms as p38 kinase inhibitors

IN Lumas, Jacques; Riedl, Bernd; Khire, Uday; Sibley, Robert N.; Hatoum-Mokdad, Holia; Monahan, Mary-katherine; Gunn, David E.; Lowinger, Timothy B.; Scott, William J.; Smith, Roger A.; Wood, Jill E.

PA Pfizer Corporation, USA

SO U.S. Pat. Appl. Publ., 39 pp., Cont.-in-part of U. S. Ser. No. 778,039.

CODEN: USXXCO

DT Patent

LA English

FAN CN 2

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	US 2002065296	A1	20020530	US 2001-838286	20010420
PRAI	US 1999-115878P	P	19990113		
	US 1999-257265	B1	19990225		
	US 1999-425229	A2	19991022		
	US 2001-778039	A2	20010207		

OS MARPAT 137:6098

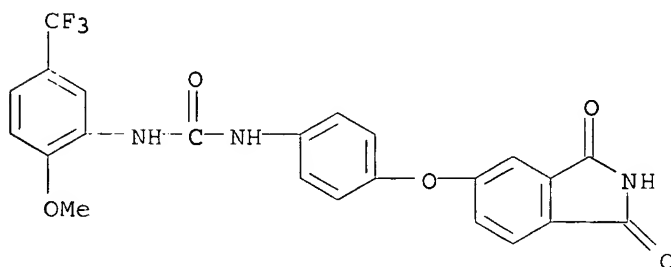
IT 274461-54-7P, N-[2-Methoxy-5-(trifluoromethyl)phenyl]-N'-[4-(1,3-dioxoisindolin-5-yl)oxy]phenyl]urea

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of heteroaryl ureas contg. nitrogen hetero-atoms as p38 kinase inhibitors)

RN 274461-54-7 CAPLUS

CN Urea, N-[4-[(2,3-dihydro-1,3-dioxo-1H-indol-5-yl)oxy]phenyl]-N'-[2-methoxy-5-(trifluoromethyl)phenyl]- (9CI) (CA INDEX NAME)



L5 ANSWER 4 OF 32 CAPLUS COPYRIGHT 2002 ACS

AB Chem. structures have been identified which allosterically modify pyruvate kinase and inhibit enzymic activity. These compds. can be used as pharmaceuticals in the treatment of a wide variety of diseases and disorders where influencing metabolic processes is beneficial, e.g. the glycolytic pathway, all pathways which use ATP as an energy source, and all pathways which involve 2,3-diphosphoglycerate related to the delivery of oxygen by modifying Hb's oxygen affinity, treatments of tumor and cancer and Alzheimer's disease. Prepn. of e.g. 2-phenylethoxy-5-formylbenzoic acid is described.

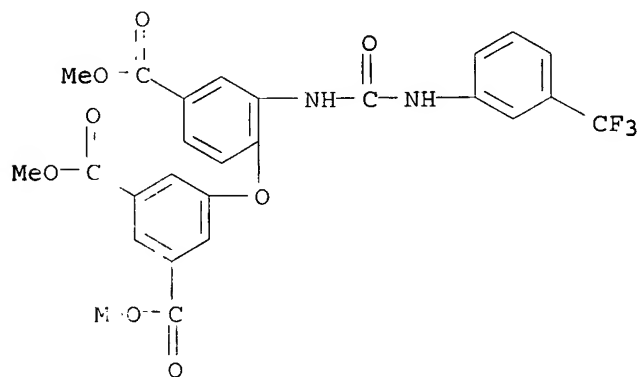
AN 01:869018 CAPLUS
 DN 06:700
 TI Allosteric inhibitors of pyruvate kinase for therapeutic use
 IN Abraham, Donald J.; Wang, Changging; Dan-Danquah, Richmond; Burnett, James C.; Joshi, Gajanan S.; Hoffman, Steven J.
 PA USA
 SO U.S. Pat. Appl. Publ., 15 pp., Cont.-in part of U.S. 6,214,879.
 CODEN: USXXCO
 DT Patent
 LA English
 FAN.CM 2

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	US 2001046997	A1	20011129	US 001-799873	20010307
	US 6214879	B1	20010410	US 098-46643	19980324
PRAI	US 1998-46643	A2	19980324		
IT	289060-07-7				

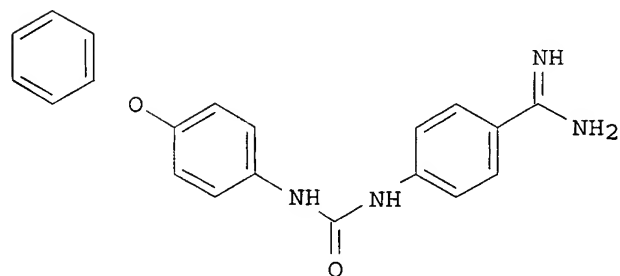
RL: BSU (Biological study, unclassified); PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
 (pyruvate kinase allosteric inhibitor for therapeutic use)

RN 289060-07-7 CAPLUS

CN 3-Benzenedicarboxylic acid, 5-[4-(methoxycarbonyl)-2-[[[3-(trifluoromethyl)phenyl]amino]carbonyl]amino]phenoxy]-, dimethyl ester
 (CA INDEX NAME)



L5 ANSWER 5 OF 32 CAPLUS COPYRIGHT 2002
 GI



I

AB Malarial parasites rely on aspartic proteases called plasmeepsins to digest Hb during the intraerythrocytic stage. Plasmeepsins from *Plasmodium falciparum* and *Plasmodium vivax* have been cloned and expressed for a variety of structural and enzymic studies. Recombinant plasmeepsins possess kinetic similarity to the native enzymes, indicating their suitability for target-based antimalarial drug development. We developed an automated assay of *P. falciparum* plasmeepsin II and *P. vivax* plasmeepsin to quickly screen compds. in the Walter Reed chem. database. A low-mol.-mass (346 Da) diphenylurea der. [WR268961 (I)] was found to inhibit plasmeepsins with a K_i of 1 to 6 μ M. This compd. appears to be selective for plasmeepsin, since it is a poor inhibitor of the human aspartic protease cathepsin D (K_i greater than 280 μ M). I inhibited the growth of *P. falciparum* strains W2 and D6, with 50% inhibitory concns. ranging from 0.03 to 0.16 μ g/mL, but is much less toxic to mammalian cells. The Walter Reed chem. database contains over 1,500 compds. with a diphenylurea core structure, 9 of which inhibit the plasmeepsins, with K_i values ranging from 0.05 to 0.68 μ M. These nine compds. show specificity for the plasmeepsins over human cathepsin D, but they are poor inhibitors of *P. falciparum* growth in vitro. Computational docking expts. indicate how diphenylurea compds. bind to the plasmeepsin active site and inhibit the enzyme.

AN 701:623551 CAPLUS

DN 135:327005

TI New class of small nonpeptidyl compounds blocks *Plasmodium falciparum* development in vitro by inhibiting plasins

AU Jiang, Suping; Prigge, Sean T.; Wei, Le Gao, Yu-E.; Hudson, Thomas H.;
Gerena, Lucia; Dame, John B.; Kyle, Der E.

CS Department of Parasitology, Division of Experimental Therapeutics, Walter Reed Army Institute of Research, Silver Spring, MD, 20910-7500, USA

SO Antimicrobial Agents and Chemotherapy, 45(9), 2577-2584
CODES: AMACCO; ISSN: 0066-4804

PB American Society for Microbiology

DT Journal

LA English

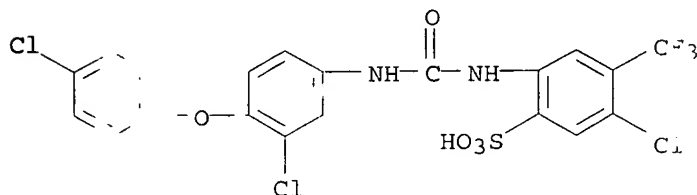
IT 447-79-0, WR 100081

RL: BAC (Biological activity or effects except adverse); BSU (Biological study, unclassified); THU (Therapeutic uses); BIOL (Biological study); USES (Uses)

(new class of small nonpeptidyl compounds blocks *Plasmodium falciparum* development in vitro by inhibiting trypsin-like proteases)

RN 447-79-0 CAPLUS

CN Ferrocenesulfonic acid, 5-chloro-2-[[[3-chloro-4-(4-chlorophenoxy)phenyl]amino]carbonyl]amino-4-(trifluoromethyl)- (9CI) (CA 76333 NAME)



RE.CN1 1 2 THERE ARE 30 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE FOLLOWING FORMAT

L5 ANSWER 6 OF 32 CAPLUS COPYRIGHT 2000 CS
GI

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

AB Title compds. [I; X = O, S; A = 1,4-C₆H₄, 1,3-C₆H₄, 1,7-naphthyl; L = H, 2,6-(CH₃)₂, 2-(CH₃)₃C, 6-(CH₃)₃C; R₁ = H, NHCO, CH₃CH₂NHCO, (CH₃)₃CNHCO, CH₃(CH₂)₅NHCO, CF₃NHCO, C₆H₅NHCO, 2-CH₃C₆H₄NHCO, 3-CH₃C₆H₄NHCO, 4-CH₃C₆H₄NHCO, 2,6-(CH₃)₂C₆H₃NHCO, 4-CH₃C₆H₄NHCO, 2,3-F₂C₆H₄NHCO; q = 0-8; m = 0-8; n = 0-8] and pharmacol. acceptable salts, which are useful as therapeutic and/or preventive agents for diabetes, hyperlipemia, arteriosclerosis, cancers, are prepd. Also, the title compd. II was prepd.

AN 2000:742094 CAPLUS

DN 133:296435

TI Preparation of amine derivatives useful as agents for diabetes, hyperlipemia, arteriosclerosis, and cancer

IN Fujita, Takashi; Wada, Kunio; Oguchi, Masaharu; Honma, Hidehito; Fujiwara, Toshihiko

PA Sankyo Company, Limited, Japan

SO Pat. Int. Appl., 208 pp.

CLASSEN: PIXXD2

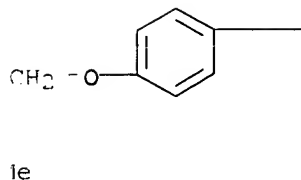
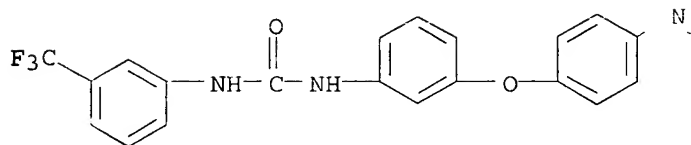
DT Patent

LA Japanese

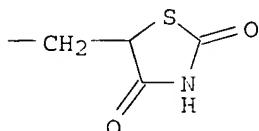
FAN.CNT

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2000061581	A1	20001019	2000-JP2216	20000406
W: AU, BR, CA, CN, CZ, HU, ID, IN, KR, MX, NO, NZ, PL, RU, TR, US, ZA				
RW: AT, BE, CH, CY, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE				
JP 2000351779	A2	20001219	2000-104702	20000406
EP 1167366	A1	20020102	2000-915362	20000406
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, NL, SE, MC, PT, IE, FI				
BR 2000009594	A	20020604	2000-9594	20000406
NO 2001004847	A	20011207	2001-4847	20011005
PRAI JP 1999-99981	A	19990407		
WO 2000-JP2216	W	20000406		
OS MARPAT 133:296435				
IT 301548-73-2P				
R: BAC (Biological activity or effect except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)				
(prepn. of amine derivs. as useful agents for diabetes, hyperlipemia, arteriosclerosis, and cancer)				
RN 301548-73-2 CAPLUS				
CN Urea, N-[3-[[2-[[4-[(2,4-dioxo-5-thiazol-5-yl)methyl]phenoxy]methyl]-1-methyl-1H-benzimidazol-6-yl]oxy]phenyl]-[3-(trifluoromethyl)phenyl]-(9CI) (CA INDEX NAME)				

PAGE 1-A

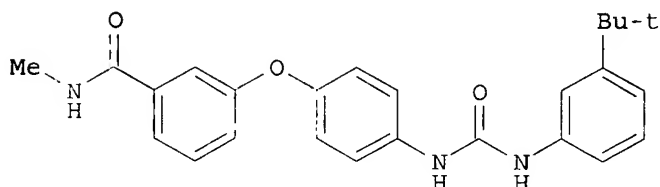


PAGE 1-B



RE.CNT 16 THERE ARE 16 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN FULL TEXT FORMAT

L5 ANSWER 7 OF 32 CAPLUS COPYRIGHT 2002 13
GI



AB This invention relates to the prepn. and use of (hetero)aryl ureas
ANHCONHB [I; A = L(ML1)q; L = 5- or 6 membered (hetero)aryl, esp. Ph or
pyridinyl; M = bridging group; L1 = (un)substituted sulfamoyl, carboxy, or
(un)substituted sulfamoyl substituent; q = 1-3; B = (un)substituted
certain (un)substituted mono- to triphenyl or heteroaryl groups] for
the treatment of raf mediated diseases, such as cancer (no data). Approx.
100 invention compds. and numerous intermediates were prepd. For
instance, 3-tert-butylaniline was coupled with
bis(trichloromethyl)carbonate to form the corresponding isocyanate, followed by addn. of
4-(3-N-methylcarbamoylphenoxy)aniline (n. given) to afford the urea
II.

AN 2000:493516 CAPLUS

DN 133:120157

TI Preparation of omega-carboxy(hetero)aryl substituted diphenyl ureas as
raf kinase inhibitors

IN Riedl, Bernd; Dumas, Jacques; Khire, Anand; Lowinger, Timothy B.; Scott,
William J.; Smith, Roger A.; Wood, Jill; Monahan, Mary-Katherine;
Natero, Reina; Renick, Joel; Sibley, Robert N.

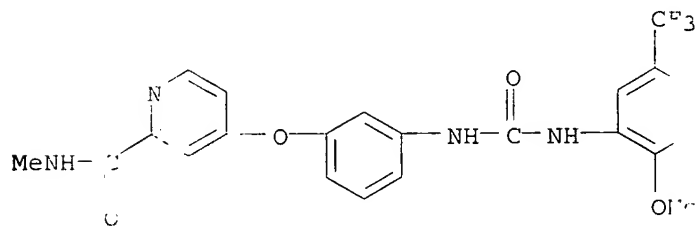
PA Bayer Corporation, USA

SO PCT Int. Appl., 120 pp.

CODEN: PIXXD2

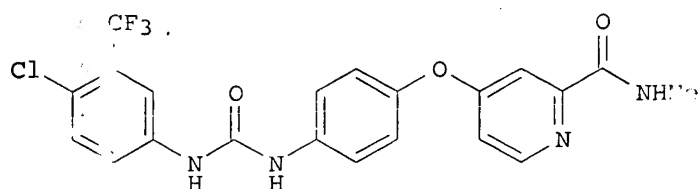
DT Patent
LA English
FAN.CNT 1

	PATENT NO.	KIND	DATE	A	ICATION NO.	DATE
PI	WO 2000042012	A1	20000720		00-US648	20000112
	W: AE, AL, AM, AT, AU, AZ, BA, BE, BR, BY, CA, CH, CN, CR, CU, CZ, DE, DK, DM, EE, ES, FI, GL, GB, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LG, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, NO, NZ, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, RW: GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, SN, TD, TG					
	EP 1140840	A1	20011010		00-903239	20000112
	R: AT, BE, CH, DE, DK, ES, FR, GB, IE, SI, LT, LV, FI, RO					
	US 2001011135	A1	20010802		773659	20010202
	US 2001011136	A1	20010802		773675	20010202
	US 2001016659	A1	20010823		773672	20010202
	US 2001027202	A1	20011004		773658	20010202
	US 2001034447	A1	20011025		773604	20010202
	US 2001003463	A	20010912		3463	20010712
	US 2002042517	A1	20020411		948915	20010910
PRAI	US 1999-115877P	P	19990113			
	US 1999-257266	A2	19990225			
	US 1999-425228	A2	19991022			
	WO 2000-US648	W	20000112			
OS	MARPAT 133:120157					
IT	284461-42-3P					
	PL: BAC (Biological activity or effect); study, unclassified); RCT (Reactant); Therapeutic use); BIOL (Biological Reactant or reagent); USES (Uses) (prepn. of .omega.-carboxy(hetero)aromatic kinase inhibitors by reacting arylamines with arylamines)					
RN	284461-42-3 CAPLUS					
CN	2-Pyridinecarboxamide, 4-[3-[[[2-methyl-4-(trifluoromethyl)phenyl]amino]carbonyl]amino]phenoxy]-N-methyl- (9CI INDEX NAME)					



RE.CNT 12 THERE ARE 12 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE TOPNAT

L5 ANSWER 8 OF 32 CAPLUS COPYRIGHT 2000
GI



AB The title compds. ADB [I; D = NHCONH: A substituted moiety of up to 40 carbon atoms of the formula L(ML1)q where L = 5-6 membered cyclic structure; L1 = substituted cyclic moiety having at least 5 members; M = bridging group having at least one atom of N, O and S; each of L and L1 contains 0-4 members of the group consisting of N, O and S); B = (un)substituted up to tricyclic aryl moiety of up to 30 carbon atoms with at least one 6-membered ring; q = 1-4; D contg. 0-4 members of the group consisting of N, O and S], useful in treating p38 mediated diseases, were prepared from the urea II which showed IC50 of 0.01-200 μ M against p38, was given. Compds. I are effective at 0.01-200 μ M against p38 (oral administration).

AN 2000:493376 CAPLUS

DN 133:120155

TI Preparation of .omega.-carboxy aryl substrates and diphenyl ureas as p38 kinase inhibitors

IN Fiedl, Bernd; Dumas, Jacques; Khire, H. S.; Nowinger, Timothy B.; Scott,
William J.; Smith, Roger A.; Wood, J. R.; Monahan, Mary-Katherine;
Matero, Reina; Renick, Joel; Sibley, J. R.

PA Eaver Corporation, USA

SO INT Int. Appl., 148 pp.

CO DEN: PIXXD2

DT Patent

LA English

FAN. CN T 2

PATENT NO.	KIND	DATE	A	CH	ON NO.	DATE
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PI	WD 2000041698	A1	20000720	US768	20000113
	W:	AE, AL, AM, AT, AU, AZ, BA, BB, BC, BD, BE, BF, BG, BH, BI, BJ, BK, BL, BM, BN, BO, BR, BS, BT, BU, BV, BW, BY, CA, CH, CN, CR, CU, CZ, DE, DK, DM, EE, ES, FI, FO, FR, GB, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LA, LB, LC, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, NO, NP, NR, NT, NU, OZ, PA, PE, PF, PG, PH, PK, PL, PM, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UY, UZ, VN, YU, ZA, ZW, AM,			

RW: GH, GM, KE, LS, MW, SD, SL, SN, ST, SZ, TD, TG, TH, TN, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, MU, NE, NG, NO, OM, PK, SI, SD, TD, TG

ED 1158985 A1 20011205 C 05597 20000113

R: AT, BE, CH, DE, DK, ES, FR, GR, IE, LI, LU, NL, SE, MC, PT,
IE, SI, LT, LV, FI, RO

PRAI US 1999-115878P P 19990113

US 1999-257265 A2 19990225

US 1999-425229 A2 19991022

WD 2000-US768 W 20000113

OS MARPAT 133:120155

IT 284461-86-5P

F.: BAC (Biological activity or effect); BSU (Biological
 study, unclassified); RCT (Reactant); THU (Therapeutic
 use); BIOL (Biological); STEP (Preparation); RACT
 (Reactant or reagent); USES (Uses)

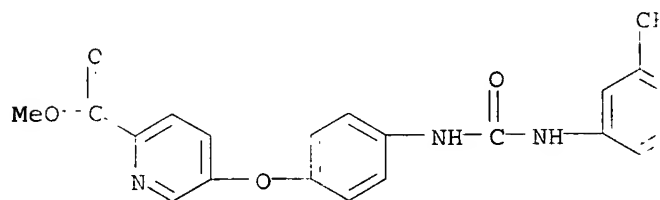
(prepn. of .omega.-carboxy aryl s s
inhibitors)

RN 134461-86-5 CAPLUS

CN 1 Pyridinecarboxylic acid, 5-[4-[[[
(trifluoromethyl)phenyl]amino]carbon
(CA INDEX NAME)

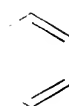
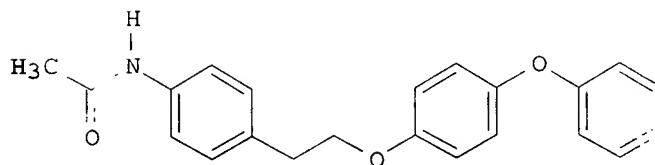
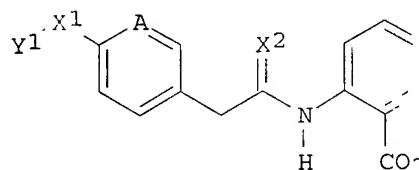
di-Ph ureas as p38 kinase

henoxy]-, methyl ester (9CI)



RE.CNT 1 THERE ARE 1 CITED REFERENCE. AP E FOR THIS RECORD
ALL CITATIONS AVAILABLE IN E T.MAT

L5 ANSWER 9 OF 32 CAPLUS COPYRIGHT 20
G1



CO₂H II

AB Title compds. [I; wherein Y1 = a group
(un)substituted-2-naphthyl; X1 is O, N, or S;
stereoisomers are prepd. and tested
therefore useful as preventive or therapeutic
agents and having cytotoxic activities useful
compd. II was prepd.

AN 0000:84754 CAPLUS

DN 132:151571

TI Preparation of anthranilic acid derivative
agents

IN Tsuchiya, Naoki; Takeuchi, Susumu; T.
Takao; Tsuruo, Takashi

represented by (un)substituted-Ph,
O or S; A = CH, N] and
agents of IgE antibody,
agents for allergic diseases
tumor agents. The title

preventive or therapeutic

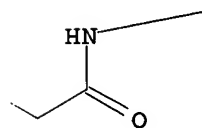
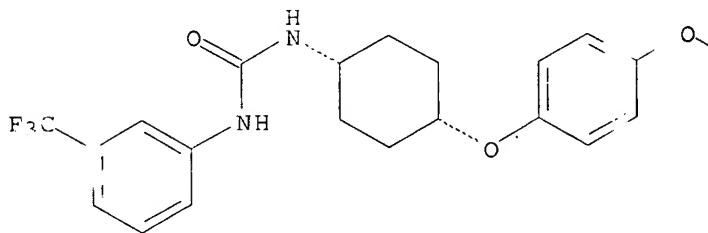
akumi; Hase, Naoki; Yamori,

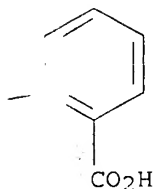
PA Teijin Limited, Japan
 SO PCT Int. Appl., 213 pp.
 CODEN: PIXXD2
 DT Patent
 LA Japanese
 FAN.CNT 1

	PATENT NO.	KIND	DATE	AP	PN NO.	DATE
PI	WO 2000005198	A1	20000203	WC	JP3969	19990723
	W:	AE, AL, AM, AT, AU, AZ, BA, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VM, YU, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM				
	RW:	GH, GM, KE, LS, MW, SD, SL, SZ, ES, FI, FR, GB, GR, IE, IT, LI, CI, CM, GA, GN, GW, ML, MR, NI, AU 9948004	A1	20000214	AU	2004 19990723
	EP 1101755	A1	20010523	EP	1522	19990723
	R:	AT, BE, CH, DE, DK, ES, FR, IE, SI, LT, LV, FI, RO				
PRAI	JP 1998-209410	A	19980724			
	JP 1998-258486	A	19980911			
	JP 1998-369808	A	19981225			
	JP 1998-369809	A	19981225			
	WO 1999-JP3969	W	19990723			
OS	MARPAT 132:151571					
IT	257606-49-8P					
	RL: SPN (Synthetic preparation); THU (Th study); PREP (Preparation); USES (Uses) (prepn. of anthranilic acid derivative agents)					
RN	257606-49-8 CAPLUS					
CN	Benzoic acid, 2-[[[4-[4-[[[cis-4-[[[2-(4-methylphenyl)amino]carbon amino]- (9CI) (CA INDEX NAME)					

Relative stereochemistry.

PAGE 1-A





RE.CNT 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RECORD

L5 ANSWER 10 OF 32 CAPLUS COPYRIGHT 2002 A
AB A method of treating a p-38 mediated disease by the
administration of BNHCONHA [A = (substituted) aryl, heteroaryl contg. 1-4
contg. 0-4 N, O, or S atoms]. Thus, 5-tert-butyl-2-(3-
tetrahydrofuranyloxy)aniline (prepn. give stirred 8 h in PhMe to give 75% N-(5-tert-
tetrahydrofuranyloxy)phenyl)-N'-(4-methylphenyl)urea. Title compds.
inhibited p38 kinase with $\text{IC}_{50} = 1-10 \text{ } \mu\text{M}$

AN 1999:421667 CAPLUS

DN 131:58659

TI Preparation of diaryl ureas as inhibitors of p38 kinase.
IN Miller, Scott; Osterhout, Martin; Dumas, Robert; Wang, Ming
Timothy Bruno; Riedl, Bernd; Scott, William; Smith, Roger A.; Wood,
Jill E.; Gunn, David; Hatoum-Mokdad, Holi; Khire, Uday; Lowinger,
Robert; Wang, Ming

PA Bayer Corporation, USA

SO PCT Int. Appl., 107 pp.

CODEN: PIXXD2

DT Patent

LA English

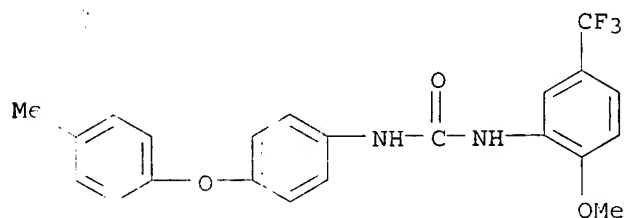
FAN.CNT 1

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WO 9932463	A1	19990701	WO	9932463	19981222
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DK, EE, ES, FI, GB, GD, GE, GH, G				HU, ID, IL, IN, IS, JP,	
KE, KG, KP, KR, KZ, LC, LK, LR, L				LU, LV, MD, MG, MK, MN,	
MW, MX, NO, NZ, PL, PT, RO, RU, S				SG, SI, SK, SL, TJ, TM,	
TR, TT, UA, UG, UZ, VN, YU, ZW, A				BY, KG, KZ, MD, RU, TJ, TM	
RW: GH, GM, KE, LS, MW, SD, SZ, UG, Z				BE, CH, CY, DE, DK, ES,	
FI, FR, GB, GR, IE, IT, LU, MC, N				SE, BF, BJ, CF, CG, CI,	
CM, GA, GN, GW, ML, MR, NE, SN, T					
CA 2315715	AA	19990701	CA	2315715	19981222
AU 9219399	A1	19990712	AU	9219399	19981222
EP 1042305	A1	20001011	EP	1042305	19981222
R: AT, BE, CH, DE, DK, ES, FR, GB, G				LI, LU, NL, SE, MC, PT,	
IE, SI, LT, LV, FI, RO					
JP 2001526276	T2	20011218	JP	2001526276	19981222
PRAI US 1997-995749	A	19971222			
WO 1998-US27265	W	19981222			
OS MARPAT 131:58659					
IT 228399-63-1P					
FI: BAC (Biological activity or effector,				t adverse); BSU (Biological	
study, unclassified); SPN (Synthetic prep				; THU (Therapeutic use);	

BIOL (Biological study); PREP (Preparation)
 (prepn. of diaryl ureas as inhibitors
 RN 228399-63-1 CAPLUS
 CN Urea, N-[2-methoxy-5-(trifluoromethyl)phenoxy]phenyl]- (9CI) (CA INDEX N

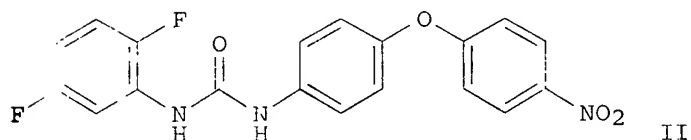
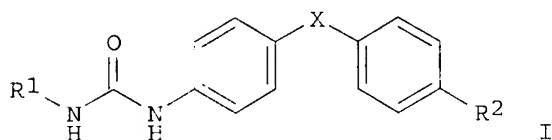
PS (Uses)
 kinase)

- [4- (4-



RE. INT 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD
 ALL CITATIONS AVAILABLE IN THE RECORD

L5 ANSWER 11 OF 32 CAPLUS COPYRIGHT 2002 A
 G



AB The invention relates to 1,3-disubstituted aryl; R2 = NO2, NH2; X = O, S], and a meta-aryl amine with isocyanates. The isocyanate reaction carried out in a solvent such as toluene, at, e.g., 80.degree.C. If a nitro group is formed, in the presence of a Pd catalyst to give an amine. The obtained 1,3-disubstituted ureas are inhibitors of the activity of the enzyme acyl-coenzyme A:cholesterol acyltransferase (ACAT), and may be used to inhibit cholesterol esterification and absorption in the intestine. For instance, reaction of 4-(4'-nitrophenoxy)phenyl isocyanate gave 76% title compd. II. The rat liver ACAT at 2 .mu.M, and 58% inhibition of ACAT in rabbit intestinal mucosa, at the same concn., both in vitro

s I [R1 = (un)substituted phenyl; R2 = NO2, NH2; X = O, S], and a meta-aryl amine with isocyanates. The isocyanate reaction carried out in a solvent such as toluene, at, e.g., 80.degree.C. If a nitro group is formed, in the presence of a Pd catalyst to give an amine. The obtained 1,3-disubstituted ureas are inhibitors of the activity of the enzyme acyl-coenzyme A:cholesterol acyltransferase (ACAT), and may be used to inhibit cholesterol esterification and absorption in the intestine. For instance, reaction of 4-(4'-nitrophenoxy)phenyl isocyanate gave 76% title compd. II. The rat liver ACAT at 2 .mu.M, and 58% inhibition of ACAT in rabbit intestinal mucosa, at the same concn., both in vitro

AN 1999:421643 CAPLUS

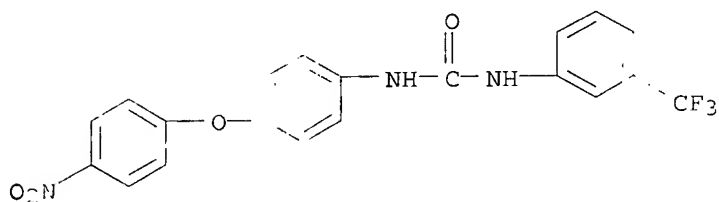
DN 131:73441

TI 1,3-Disubstituted ureas useful as ACAT inhibitors, and method for their preparation

IN Oremus, Vladimir; Smahovsky, Vendelin; Falcetta, Viera; Kakalik, Ivan;

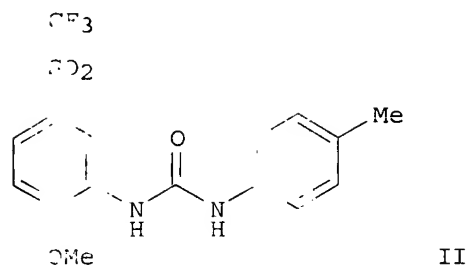
Schmidtova, Ludmila; Zemanek, Marian
 PA Slovako-Farma, A.S., Slovakia
 SO PCR Int. Appl., 33 pp.
 CODEN: PIXXD2
 DT Patent
 LA English
 FA .CNT 1

	PATENT NO.	KIND	DATE	APP	ON NO.	DATE
PI	WO 9932437	A1	19990701	WO	K19	19981216
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	KG, KP, KR, KZ, LC, LK, LR, LS, I				J, LV, MD, MG, MK, MN, MW,	
	MX, NO, NZ, PL, PT, RO, RU, SD, S				3, SI, SK, SL, TJ, TM, TR,	
	TT, UA, UG, US, UZ, VN, YU, ZW, A				7, BY, KG, KZ, MD, RU, TJ, TM	
	RW: GH, GM, KE, LS, MW, SD, SZ, UG, Z				BE, CH, CY, DE, DK, ES,	
	FI, FR, GB, GR, IE, IT, LU, MC, N				BE, BF, BJ, CF, CG, CI,	
	CM, GA, GN, GW, ML, MR, NE, SN, T					
	AU 9916976	A1	19990712	AU	6976	19981216
	EP 1042278	A1	20001011	EP	61715	19981216
	R: AT, BE, CH, DE, DK, ES, FR, GB, G				LI, LU, NL, SE, PT, IE,	
	SI, FI, RO					
	JP 2001526259	T2	20011218	JP	525374	19981216
PRA	SK 1997-1751	A	19971219			
	WO 1998-SK19	W	19981216			
OS	MARPAT 131.73441					
IT	223544-40-9P					
	RL: BAC (Biological activity or effector,				adverse); BSU (Biological	
	study, unclassified); SPN (Synthetic prep); THU (Therapeutic use);	
	BIOL (Biological study); PREP (Preparatic				US (Uses)	
	(prepn. of 1,3-disubstituted ureas as				inhibitors)	
RN	228544-40-0 CAPLUS					
CN	Urea, N-[4-(4-nitrophenoxy)phenyl]-N'-[3-				loromethyl)phenyl]- (9CI)	
	(CA INDEX NAME)					



RE.CNT 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD
 AND CITATIONS AVAILABLE IN THE P. 7 MAT

L5 ANSWER 12 OF 32 CAPLUS COPYRIGHT 2002 7
 GI



AB The invention relates to the use of a group of aryl ureas ANHCONHB [I; A = certain (un)substituted Ph, pyridinyl, or 2-yl groups; B = certain (un)substituted mono- to tricyclic aryl or heteroaryl groups] in treating raf-mediated diseases, and pharmaceutical compositions for use in such therapy. A subset of I are novel and are claimed per se. Numerous intermediates were prepared. For instance, reaction of 2-methoxy-5-(trifluoromethylsulfonyl)aniline in EtOAc gave title compd. II. In an in vitro kinase assay, all compds. displayed IC50 values between 1 nM and 10 μM.

AN 1999:421642 CAPLUS

DN 131:58658

TI Inhibition of raf kinase using symmetrical and unsymmetrical substituted diphenyl ureas

IN Miller, Scott; Osterhout, Martin; Dumas, J.; Khire, Uday; Lowinger, Timothy Bruno; Riedl, Bernd; Scott, William Smith, Roger A.; Wood, Jill E.; Gunn, David; Rodriguez, Mareli; Wang

PA Bayer Corporation, USA

SO PCT Int. Appl., 89 pp.

COEN: PIX02

DT Patent

LA English

FAI.CNT 1

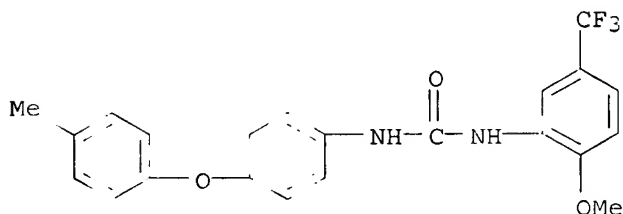
PATENT NO.	KIND	DATE	APPL. NO.	FILED NO.	DATE
WO 9932436	A1	19990701	WO 1	96081	19981222
W: AL, AM, AT, AU, AZ, BA, BB, BG, BF, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SI, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, UZ, VN, YU, ZW, AM, BY, KG, KZ, MD, RU, TJ, TM, RW: GH, GM, KE, LS, MW, SD, SE, UG, ZW, FI, FR, GB, GR, IE, IT, LU, MC, NL, CM, GA, GN, GW, ML, MR, NE, SN, TD					
CA 2315646	AA	19990701	CA 1	2315646	19981222
AU 9919054	A1	19990712	AU 1	99054	19981222
EP 1049664	A1	20001108	EP 1	103809	19981222
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IF, IL, LT, LV, FI, RO					
JP 2001526258	T2	20011218	JP 1	25373	19981222
BR 1314375	A	20020521	BR 1	4375	19981222
NO 2000003230	A	20000821	NO 2	230	20000621
PRAI US 1997-990344	A	19971222			
WO 1998-US6081	W	19981222			
OS MARPAT 131-58658					
IT 228399-63-1P					

PL: BAC (Biological activity or effector, adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use);

BIOL (Biological study); PREP (Preparation
(prepn. of sym. and unsym. substituted
effects on tumors mediated by r f kinas
RN 228399-63- CAPLUS
CN Urea, N-[2 methoxy-5-(trifluoromethyl) pher
methylphenoxy'phenyl]- (9CI) (CA INDEX N.

SES (Uses)
areas with inhibitory

4-(4-



RECENT 3 THERE ARE 3 CITED REFERENCES AVAIL
AL CITATIONS AVAILABLE IN THE RE

FOR THIS RECORD
AT

L5 ANSWER 13 ON 32 CAPLUS COPYRIGHT 2002 AC
AB Sodium 2-[2-[3-[4-chloro-3-(trifluoromethyl)
(trifluoromethyl)phenoxy]-4,5-dichlorobenz
steps from 3,4-dichlorophenol and 4-chloro
prepd. were sodium 2-[2-[3-[3,5-bis(triflu
(trifluoromethyl)phenoxy]-5-(1,1-dimethyl
sodium 2-[2-[3-[4-chloro-3-(trifluoromethyl
(trifluoromethyl)phenoxy]-5-(1,1-dimethyl
edema induced in the mouse by 12-O-tetradec
mg/ear topically, 2-[2-[3-[4-chloro-3-(tri
(trifluoromethyl)phenoxy]-5-(1,1-dimethyl
exhibited an ED50 of 0.32 mg/ear and 2-[2-
bis(trifluoromethyl)phenyl]amino]carbonyl]
(trifluoromethyl)phenoxy]-5-(1,1-dimethyl
exhibited an ED50 of 0.87 mg/ear.

ureido]-4-
onate was prepd. in 5
benzotrifluoride. Also
yl)phenyl]ureido]-4-
benzenesulfonate and
ureido]-4-
benzenesulfonate. For ear
phorbol 13-acetate at 50
omethyl)phenyl]ureido]-4-
benzenesulfonic acid
4-
benzenesulfonic acid

AN 1999:384011 APLUS
DN 131:44661
TI Anti-inflammatory compounds
IN Dixon, James Scott; Hall, Ralph Floyd; May
H., III; Mayer, Ruth Judik; Winkler, James
PA Smithkline Beecham Corporation, USA The J
SC U.S., 17 pp., Cont.-in-part of U.S. 5,470,
CODEN: USXXAM
DT Patent
LA English
FAM.CIT 2

Lisa Ann; Chilton, Floyd
Hopkins University

PATENT NO.	KIND	DATE	APP
US 5912270	A	19990615	US
US 5470882	A	19951128	US
WO 9533712	A1	19951214	WO
W: JP, US			
RW: AT, BE, CH, DE, DK, ES, FR, GB, GR			
PF I US 1994-25271		19940602	
WO 1995-US657		19950602	
OS MARPAT 131:44 61			
IT 447-64-3P			
E: BAC (Biological activity or effector,			

CN NO.	DATE
57650	19961122
57716	19940602
57677	19950602
T, LU, MC, NL, PT, SE	
adverse); BSU (Biological	

study, unclassified); SPN (Synthesis Preparation); PREP (Preparation)

(prepn. of antiinflammatory ureidophenyl

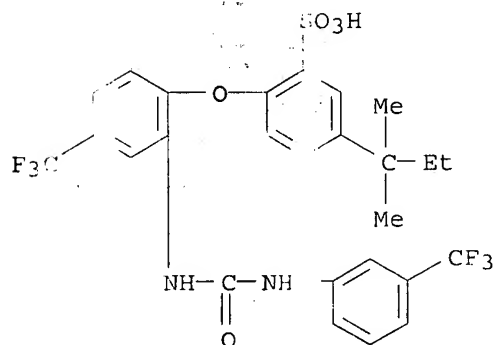
RN 447-64-3 CAPLUS

CN Benzenesulfonic acid, 5-(1,1-dimethylpropyl)-[[[3-(trifluoromethyl)phenyl]amino]carbonyl]-
INDEX NAME)

; BIOL (Biological

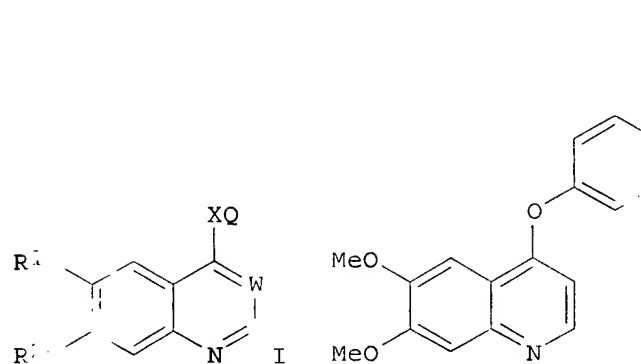
benzenesulfonates)

5-(trifluoromethyl)-2-
[3-(trifluoromethyl)phenoxy]- (9CI) (CA



RE.CN 28 THERE ARE 28 CITED REFERENCE AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE FILE

L5 ANSWER 14 OF 2 CAPLUS COPYRIGHT 1997
G1



AB The title compds. I [R1 and R2 represent H or Me; R1 and R2 together form C1 to C3 alkylene; XQ represents CH or N; and W represents substituted heteroaryl] are prepd. I inhibit platelet-derived growth factor-induced autophosphorylation and are useful in the treatment of cancer, arthritis, etc. The title compd. II (prepn. given) (1 day) increased the survival of mice with tumor cells by 130%

AI 1997:414195 APLUS

DN 127:34137

TI Preparation of quinoline and quinoxaline

R1, R2 = C1-4 alkyl, or R1 and R2 together form C1-4 alkylene; XQ represents O, S or CH2; W represents substituted heteroaryl] are prepd. I inhibit platelet-derived growth factor receptor
treatment of cancer, arthritis, etc.
100 mg/kg i.p. once daily for 14 days
implanted leukemic P388

has inhibiting

platelet-derived growth factor receptor
IN Kubo, Kazuo; Ohyama, Shinichi; Shimizu, T.
Kato, Shinichiro; Murooka, Hideko; Koga,
PA Mirin Beer Kabushiki Kaisha, Japan; K.
Shimizu, Toshiyuki; Nishitoba, Tsunosh
SC ICT Int. Appl., 243 pp.
CODEN: PIXXD2
DT Patent
LA Japanese
FAM.CNT 1

k ; Nishitoba, Tsuyoshi;
 orihiko; et al.
 ; Ohyama, Shinichi;
 Shinichiro

	PATENT NO.	KIND	DATE	APT
PI	WO 9717329	A1	19970515	WO
	W: AL, AM, AT, AU, AZ, BA, BB, BE, BG, BR, CA, CH, CN, DE, DK, EE, ES, FI, GB, GE, GR, HU, IL, IN, JP, KE, KR, KZ, LG, LU, LV, MD, MG, MK, MN, MU, MY, NL, NO, NZ, OM, OS, PA, PE, PG, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, TH, TJ, TM, TR, TT, UA, UG, UZ, VN, YU, ZA, ZM, ZW			
	RW: KE, LS, MW, SD, SZ, UG, AZ, BA, CH, CN, DE, DK, EE, ES, FI, GB, GE, GR, HU, IL, IN, JP, KR, KZ, MD, RU, TJ, TM, UA, UG, UZ, VN, YU, ZA, ZM, ZW			
	MR, NI, SN, TD, TG			
	AU 9673400	A1	19970529	AU
	EP 860433	A1	19980826	EP
	R: CH, DE, FR, GB, LI			
	US 6143764	A	20001107	US
PFPI	EP 1995-313555	A	19951107	
	EP 1996-62121	A	19960223	
	WO 1996-JP3222	W	19961105	

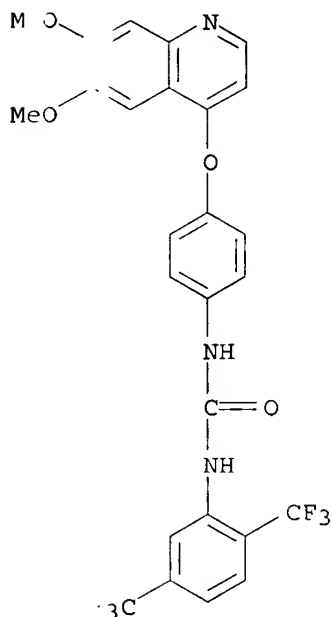
LOF NO.	DATE
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PK, NO, NZ, PL, PT, RO,	
TA, UG, US, UZ, VN, AM,	
K, ES, FI, FR, GB, GR,	
CG, CI, CM, GA, GN, ML,	
73490	19961105
935541	19961105
860	19980506

OS MARPAT 127:34 37
IT 190727-78-7P
RL: BAC (Biological activity or effect on study, unclassified); SPN (Synthetic and BIOL (Biological study); PREP (Preparation (prepn. of quinoline and quinazoline, platelet-derived growth factor receptor)
RM 190727-78-7 CAPLUS
CI Area, N-[2,5-bis(trifluoromethyl)-phenylquinolinyl]oxy]phenyl]- (9CI) (C, IN)

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pt adverse); BSU (Biological
on); THU (Therapeutic use);
SES (Uses)
    inhibiting
    phosphorylation)
[ 5,7-dimethoxy-4-

```



L5 ANSWER 15 OF 22 CAPLUS COPYRIGHT 2001
 AB CoA-independent transacylase (CoA-IT) inhibitors for inhibiting or reducing cell proliferation or inducing apoptosis, but the I is for treatment of other CoA-IT-mediated diseases. II inhibited CoA-IT at apoptosis-inducing activity. The specific is also described.

AN 1997:207756 CAPLUS

DI 126:195233

T1 Compounds for inhibition of CoA-independent transacylase, induction of apoptosis, treating CoA-independent transacylase-dependent diseases, and inhibiting cell proliferation

IN Winkler, James David; Chilton, Floyd III

PA Smithkline Beecham Corporation, USA; Winkler, James David; Chilton, Floyd III

SO PCT Int. Appl., 34 pp.

CODEN: PIXXD2

DT Patent

LA English

FAN.CNT 1

PATENT NO.	KIND	DATE	PPU	IOF NO.	DATE
WO 9704765	A1	19970213	O	US 5257	19960724
W: JP, U					
RJ: AT, B, CH, DE, DK, ES, FI, FR					
TP 841910	A1	19980520		501	19960724
I: BE, CA, DE, ES, FR, GB, IT, LI					
JP 11511130	T2	19990928		752	19960724
PAI US 1995-2239P	P	19950725			
WO 1996-US1225	W	19960724			

s are disclosed for human or mammal. Compds. osis exclude holine (I) or alkyl s are disclosed for tron. of e.g. di-Et y heptanephosphonate (II) o 9 .mu.M; II also showed bition of CoA-IT by I is

ansacylase, induction of se-dependent diseases, and

s' University; Winkler,

IT 173730-67-1P

-BAC (Biological activity or effect study, unclassified); SPN (Synthetic preparation); BIO (Biological study); PREP (Preparation of compounds for inhibition of CoA-dependent apoptosis, treating CoA-independent transcription inhibiting cell proliferation, and compounds

```

at adverse); BSU (Biological
on): THU (Therapeutic use);
RES (Uses)

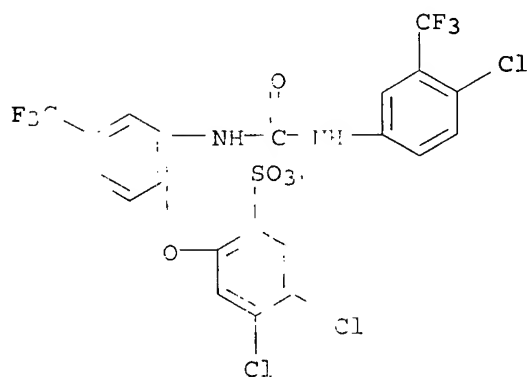
```

transacylase, induction of
 -ase-dependent diseases and
 rep .)

RI 173730-67-1 C/ PLUS

C Benzene-sulfonic acid, 4,5-dichloro-2- -[[[trifluoromethyl]phenyl]amino]carbonyl aminomorpholinium salt (9CI) (CA INDEX NAME)

chloro-3-
- (trifluoromethyl)phenoxy] -

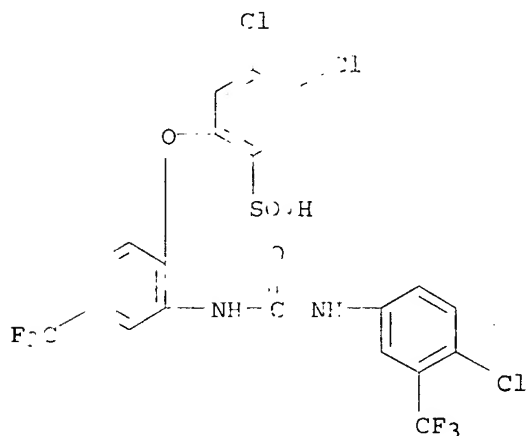


● Na

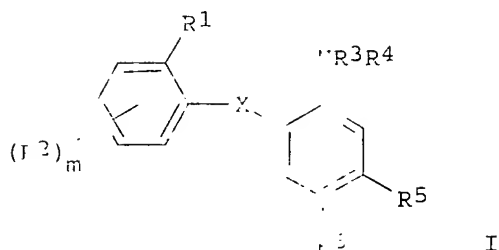
L5 ANSWER 26 OF 37 CAPLUS COPYRIGHT2006 ACS
AB ET-18-O-CH3 (1)-octadecyl-2-O-methyl n-gl
antiproliferative agent, blocking the growth
and in vivo. However, there is controversy
leading to its antiproliferative effects.
(CoA-IT) is an enzyme that remodels a cholic
phospholipid donor and acceptor molecules. In a
ET-18-O-CH3 was a potent inhibitor of CoA-IT
kinetic anal. revealed that its inhibition
lyso-phospholipid substrate. The goal of this
the connection between inhibition of CoA-IT
using several structurally distinct inhibitors
other inhibitors of CoA-IT were found to inhibit
thymidine incorporation into the DNA, as well as
humor HL-60 monocytic leukemia cells. The
by ET-18-O-CH3 appeared to be different from
necrosis factor; the former failed to activate
necrosis factor did. Closer examination of the
model revealed that compounds that were true
inhibitors, but lacked CoA-IT inhibitory activity
apoptosis. In addition, compounds that inhibit
participate in arachidonic acid metabolism
phospholipase A2, did not induce apoptosis.
demonstrate that inhibition of CoA-IT can block
proliferation and the induction of apoptosis.

ro-3-phosphocholine) is an
f cancer cells both in vitro
garding the mechanism
-independent transacylase
te between specific
iet of mammalian cells.
EC50, 0.5 .mu.M), and
competitive with the
current study was to explore
d antiproliferative effects
of CoA-IT. ET-18-O-CH3 and
it cell proliferation and
as to induce apoptosis in
hanism of apoptosis induced
hat induced by tumor
NF .kappa.B, whereas tumor
emarcol. of apoptosis in this
all related to CoA-IT
it, also failed to induce
other enzymes that
xygenase, 5-lipoxygenase and
aken together, these results
inked to blockade of
n HL-60 cells.

DN 26:166148
 TI Inhibitors of coenzyme A-independent trans-ase induce apoptosis in
 Human HL-60 cells
 AU Linker, James E.; Eris, Tamer; Sung, Hui ; Cilibot-Fletcher, Marie;
 Mayer, Ruth J.; Surette, Marc E.; Chilton, yd
 CE Dep. Immunopharmacol. Med. Chem., Smit Klir eed am Pharmaceuticals, King
 of Prussia, PA, USA
 SO Journal of Pharmacology and Experimental Th pectics (1996), 279(2),
 956-966
 CODEN: JPETAR; ISSN: 0022-3565
 PE Williams & Wilkins
 DT Journal
 LA English
 IF 162793-63-7, Skf 45905
 AB BAC (Biological activity on effector, pt course); BSU (Biological
 study, unclassified); BIOL (Biological stu induce apoptosis in human
 (inhibitors of CoA-independent trans acyl
 HL-60 cells)
 RN 162793-63-7 CAPLUS
 CM Benzenesulfonic acid, 4,5-dichloro-2- [[chloro-3-
 (trifluoromethyl)phenyl]amino]carbonylamin -4-(trifluoromethyl)phenoxy]-
 (DCI) (CA INDEX NAME)



LE ANSWER 17 0 32 CAPLUS COPYRIGHT 10 AC
 G



AI The invention relates to the novel compounds of I
 [R1 = SO3H, S(O)n-C1-4 alkyl; n = 0-2; R2 = (substituted) C1-8
 alkyl, C1-8 alkoxy; m = 1, 2; P = C(C(R7, ()R7, R4, R8, R9 = H, C1-4
 alkyl; R5 = H, halo, CF3, Me, (CH2)tC(O)2R8 CH2 tOH; t = 0-2; R6 = H,
 halo; R7 = (substituted) aryl, (substituted) aryl C1-2 alkyl,
 (substituted) C1-8 alkyl, NR9R10; R10 = (substituted) aryl, (substituted)
 aryl-C1-2 alkyl, (substituted) C1-8 alkyl, R9NR10 form 5- to 7-membered
 (un)satd. ring with optional addl. hetero n of O/N or S; X = O, S; with
 provisions] and pharmaceutically acceptable salts thereof. The invention
 also relates to a method of treating a condition of inflammation in a mammal
 in need thereof, which comprises administering to said mammal an effective
 amt. of a compd. or compn. of I. Prep. of selected compds. of the
 invention is described. Compds. of the invention demonstrated
 phospholipase A2 inhibition, generally at submicromolar levels.

AI 1996:137692 CARLUS

DI 124:165248

TI Aryl antiinflammatory compounds, their preparation, and their activity

IN Adams, Jerr; McCoy; Hall, Ralph Floyd

PA SmithKline Beecham Corp., USA

SC ACT Int. Appl. 16 pp.

CO DEN: PIXX02

DI Patent

LA English

FAN. CTT 1

PATENT NO.	KIND	DATE	FILE NO.	DATE
WO 9533458	A1	19951214	US6951	19950602
RW: AT, BF, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE				

PRAT US 1994-252 8 19940602

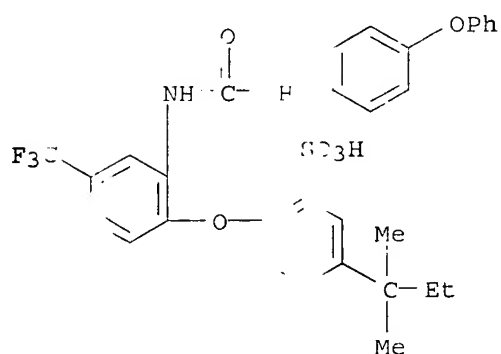
OS PARPAT 124:165248

IT 174083-25-1P

AB: BAC (Biological activity or effect); SPN (Synthetic preparation); THU (Therapeutic use); IOL (Industrial study); PREP (Preparation); USES (Uses)
 (aryl antiinflammatory compd. prepn and activity)

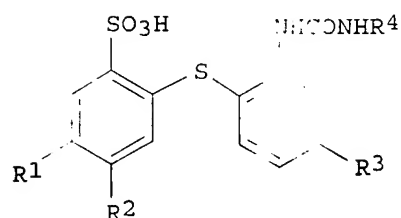
RN 174083-25-1 CARLUS

CN Benzenesulfonic acid, 5-(1,1-diethyl-2-[[[4-(bromomethyl)phenoxy]-, monosodium salt (9CI) (CA INDEX NAME

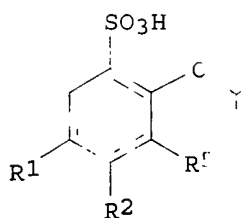


● Na

L5 ANSWER 18 CAPLUS COPYR 197200 A
GI



I



II

AB Pharmaceutical compns. are disclosed which contain I (R1 = Cl; R2 = H, Cl, R3 = Cl, CF3; R4 = Ph substituted at 1-2 positions with Cl or CF3; when R1 = Cl, C((CH3)2CH2CH3; R2 = substituted at 1-2 positions with Cl or CF3, or disubstituted Ph substituted once by Cl or CF3 and once by 3-chlorophenoxy or 4-chlorophenoxy; with pharmaceutically acceptable diluent or carrier) and a method for treating or reducing inflammation in a mammal by administering an effective amt. of a compd. or compns. of selected compds. of the invention as defined herein.

AN 1996:13285 CAPLUS

DN 124:165243

TI Anti-inflammatory benzenesulfonic acid derivatives and their activity

IN Dixon, James G.; Hall, Ralph F.; Marshall, III; Mayer, Fred J.; Winkler, James D.

PA SmithKline Beecham Corp., USA

SO U.S., 16 pp.

CODEN: USXXAM

DT Patent

LA English

FAN.CNT 2

ONHR 1

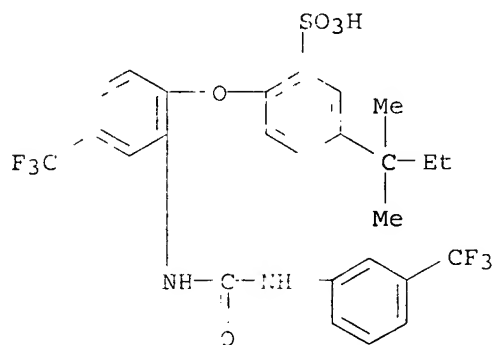
R2

contain I (R1 = Cl; R2 = H, Cl, R3 = Cl, CF3; when R1 = Cl, C((CH3)2CH2CH3; R2 = substituted at 1-2 positions with Cl or CF3, or disubstituted Ph substituted once by Cl or CF3 and once by 3-chlorophenoxy or 4-chlorophenoxy; with pharmaceutically acceptable diluent or carrier) and a method for treating or reducing inflammation in a mammal by administering an effective amt. of a compd. or compns. of selected compds. of the invention as defined herein.

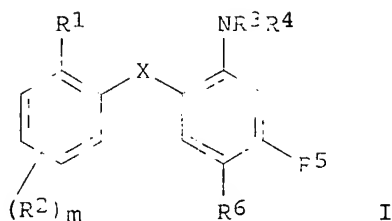
tives, their preparation, and

sa A.; Chilton, Floyd H.,

	PATENT NO.	KIND	DATE	APPL. NO.	DATE
PI	US 5470882	A	19951128	U	4-25-11 19940602
	WO 9533712	A1	19951214	V	5-US6577 19950602
	W: JP, US				
	RW: AT, BE, CH, DE, DK, ES, FR, GB,				E, IT, LU, MC, NL, PT, SE
	FP 765305	A1	19970407	EP	-922898 19950602
	EP 765305	B1	19991215		
	R: BE, CH, DE, FR, GB, IT, LI, NL				
	JP 10506092	T2	19980615	JP	500061 19950602
	US 5912270	A	19990615	US	-720050 19961122
PRAI	US 1994-252710		19940602		
	WO 1995-US6677		19950602		
OS	MARPAT 124:168243				
IT	447-64-3				
	RL: BAC (Biological activity or effector, (Therapeutic use); BIOL (Biological study; (anti-inflammatory benzenesulfonic acid their activity)				ept adverse); THU SES (Uses) rives., their prepn., and
RN	147-64-3 CAPLUS				
CN	Benzenesulfonic acid, 5-(1,1-dimethylpropyl)- [[[3-(trifluoromethyl)phenyl]amino]carbamoyl- INDEX NAME)				2-[(trifluoromethyl)-2- ami phenoxy]- (9CI) (CA



L5 ANSWER 19 OF 22 CAPLUS COPYRIGHT 2002
GI



AB This invention relates to the novel comp and pharmaceutical compns. of
formula I wherein R1 is (CH2)nH or (CH2 R8 ; n is 0 or an integer

having a value of 1; X is oxygen or sulfur, optionally substituted C1-8 alkyl, or C1-4 alkyl having a value of 1 or 2; R3 is C(O)R7; R4 is hydrogen, halogen, CF3, CH3, (CH2)tCO2R9; t is an integer having a value of 1 or 2; R6 is hydrogen, halogen, CF3, CH3, (CH2)tCO2R9; R8 is hydrogen or C1-4 alkyl; R9 is hydrogen, optionally substituted aryl, or C1-4 alkyl, optionally substituted C1-8 alkyl, or nitrogen to which they are attached form an unsatd. ring which may optionally comprise a heteroatom selected from O/N or sulfur; or a pharmaceutically acceptable salt thereof, which comprises an effective amount of a compd. or compn. of 2-[2-[3-(4-bromophenyl)ureido]-4-(trifluoromethyl)phenoxy]benzoic acid which was hydrogenated over 10% Pd/C to 2-[2-[3-(4-bromophenyl)ureido]-4-(trifluoromethyl)phenoxy]benzoic acid which inhibited PLA2 and CoA-IT at 500 nM or

AN 1995:838690 CAPLUS
 DN 124:8418
 TI Antiinflammatory (ureidophenoxy)benzoic acid inhibitors of phospholipase A2 and CoA-inhibitors
 IN Adams, Jerry L.; Hall, Ralph F.; Seibel, William R.
 PA SmithKline Beecham Corp., USA
 SO U.S., 17 pp.
 CODEN: USMAA1
 DT Patent
 LA English
 FAN.CNT 1

PATENT NO.	KIND	DATE	APPL. NO.	FILED DATE
US 5447957	A	1995090	US 5447957	19940602
WO 9533460	A1	19951214	WO 9533460	19950602

W: EP, US

RW: AT, AU, BE, CH, DE, DK, ES, FR, GB, G

PRAI US 1994-03-14 19940602

OS 124:8418

IT 17.103-10-9P

TL: BAC (Biological activity or effector, preparation); THU (Therapeutic use); BIOL (Biological preparation); USES (Uses)

(antiinflammatory (ureidophenoxy)benzoic acid inhibitors of phospholipase A2 and CoA-inhibitors)

RN 17.103-10-9 CAPLUS

CN 2-[2-[3-(4-bromophenyl)ureido]-4-(trifluoromethyl)phenoxy]benzoic acid (CA INDEX 124:8418)

R2 is hydrogen, halogen, alkoxyl, or C1-4 alkyl; R5 is hydrogen, halogen, CF3, CH3, (CH2)tCO2R9; t is 0 or an integer having a value of 1 or 2; R6 is hydrogen or halogen; R7 is NR9R10 or C1-4 alkyl; R10 is hydrogen, optionally substituted aryl, or C1-4 alkyl, optionally substituted C1-8 alkyl, or nitrogen to which they are attached form an unsatd. ring which may optionally comprise a heteroatom selected from O/N or sulfur; or a pharmaceutically acceptable salt thereof. This compound is useful for reducing inflammation in a mammal by administering to said mammal an effective amount of 2-[2-[3-(4-bromophenyl)ureido]-4-(trifluoromethyl)phenoxy]benzoic acid which

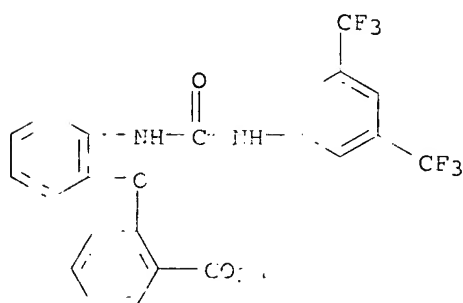
is an effective amount of 2-[2-[3-(4-bromophenyl)ureido]-4-(trifluoromethyl)phenoxy]benzoic acid which

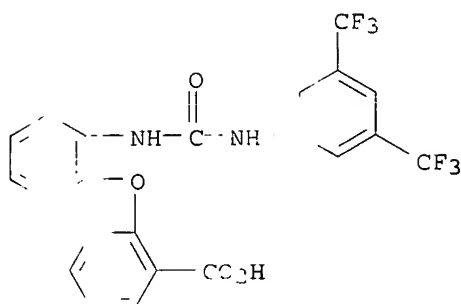
AT, AU, BE, CH, DE, DK, ES, FR, GB, G

pt adverse); SPN (Synthetic preparation); PREP (Preparation)

ic (antiinflammatory (ureidophenoxy)benzoic acid inhibitors of phospholipase A2 and CoA-inhibitors)

2-[2-[3-(4-bromophenyl)ureido]-4-(trifluoromethyl)phenoxy]benzoic acid





L5 ANSWER 20 OF 32 CAPLUS COPYRIGHT 2002 AC
 AB The enzyme CoA-independent transacylase (C
 mediate the movement of arachidonate between
 subclasses, and we have shown that two inh
 and SK&F 45905) block this movement. In t
 inhibitors to further characterize the rol
 lipid metabolism. SK&F 98625 (1-ethyl-7-(3,4
 imidazol-1-yl)heptane-phosphonate) and SK&
 (trifluoromethylphenyl)ureido-2,2,2-trifluorom
 dichlorobenzene-sulfonic acid inhibited Co
 .mu.M and 6 .mu.M, resp.). Another compd.
 cyclooxygenase, 14-kDa PLA2, and acetyltrans
 below 20 .mu.M. However, SK&F 45905 inhib
 = 3 .mu.M), and both compds. inhibited 5-l
 values of 1-4 .mu.M). In platelet-stimul
 SK&F 45905 blocked the liberation of arach
 which suggests that the movement of arachi
 phospholipid pools is a prerequisite for r
 inhibits the prodn. of platelet activatin
 neutrophils and antigen-stimulated mast ce
 platelet-activating factor and arachidonic
 by an inhibitor of 5-lipoxygenase, zileuton.
 primary mode of action of SK&F 98625 and S
 CoA-IT. SK&F 98625 and SK&F 45905 were ab
 prodn. in peritoneal inflammatory cells and t
 ears of phorbol ester-challenge mice. Tha
 that blockade of CoA-IT, which leads to ir
 remodeling between phospholipids, results
 of platelet activating factor prodn., arachid
 formation of eicosanoid products.

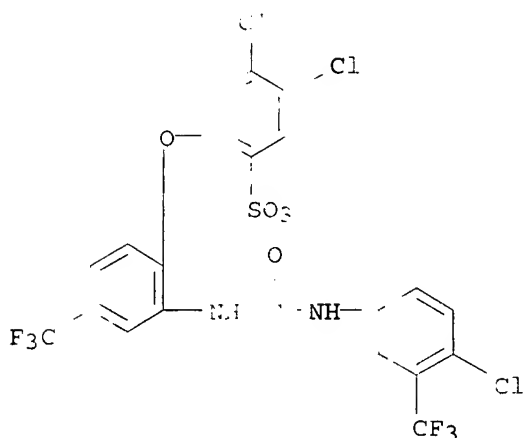
AN 1993:92:639 CAPLUS
 DN 123:175-183
 TT Effects of CoA-independent transacylase in
 lipid inflammatory mediators
 AU Winkler, James D.; Fonteh, A. F.; N.; Sung
 Nixon, Andrew B.; Chabot-Fletcher, Marie;
 A.; Chilton, Lloyd H.
 CS D. J. Pharmacol., SmithKline Beecham Pharm.
 SO J. Pharm. Mol. Exp. Ther. (1995) 273(3), 1
 J. Pharm. Mol. Exp. Ther. (1995) 273(3), 1
 DT Journal
 LA English
 IT 162793-62-7, SKF 45905
 RL BAC Biological activity of factor,
 (Biological study,

IT) has been proposed to
 specific phospholipid
 CoA-IT (SK&F 98625
 we use these
 in the prodn. of
 2-oxo-2,3-dihydro-
 [2-[3-(4-chloro-3-
 -oxy)-4,5-
 (IC50 values of 9
 effect on
 activities at concns.
 185 nM PLA2 activity (IC50
 cyclooxygenase activity (IC50
 d. In cells, SK&F 98625 and
 from phospholipids,
 specific
 both compds. also
 ionophore-stimulated
 inhibition of
 release was not mimicked
 which indicates that the
 45905 via inhibition of
 prostaglandin
 of inflammation in
 these results show
 arachidonate
 of
 release and the

the production of
 teravi, Javid D.;
 Marshall, Lisa
 PA, USA

BIOL

(effects of CoA-independent transacylase on the prodn. of lipid infl ammatory mediators)
 P 2793-07 APUS
 C Benzene sulfonic acid, 4,5-dichloro-2-[2-[(trifluoromethyl)phenyl]amino]ethylammonium salt (9CI) (CAS EX 17ME)



L5 ANSWER 21 CF 02 CAPLUS COPYRIT 2002 AC
 AB The enzyme CoA-independent transacylase (CoA-IT) is proposed to mediate the movement of arachidonate between phospholipid subclasses and influence the formation of arachidonic acid, a precursor of platelet-activating factor. To substantiate this hypothesis, the authors have developed two structurally distinct inhibitors, SK&F 98625 [diethyl 2-[3,4,5-trimethylimidazole-1-yl]heptane phosphonate] and SK&F 45905 [(trifluoromethyl)phenyl]ureido]-2-[(trifluoromethyl)phenyl]ureido]-4,5-dichlorobenzene sulfonic acid]. These compounds inhibit the capacity to block microsomal CoA-IT activity, the transacylation of 1-alkyl-2-linoleoyl-sn-glycerol-3-phosphate transfer of [¹⁴C]arachidonate from 1-acyl-lysophosphatidylcholine (lyso-PE). Both SK&F 98625 and SK&F 45905 (10⁻⁶-10⁻¹⁹ M) in these two assays. In contrast, these compounds had little or no effect on other lipid-modifying enzymes. CoA-dependent acyltransferase or acetyltransferase revealed that both SK&F 98625 and SK&F 45905 inhibit the enzyme and prevented the acylation of lysophosphatidylcholine in a competitive manner. In intact human neutrophils, both compounds completely blocked the movement of [¹⁴C]arachidonate from 1-alkyl-2-linoleoyl-sn-glycerol-3-phosphate (GPE). In contrast, these compounds did not alter the incorporation of free arachidonate into phospholipids in intact cells. This is the first report to demonstrate the importance of CoA-IT in arachidonate metabolism. These results provide further evidence that CoA-IT is a key enzyme in the large pool of -etl-lysophospholipids in human neutrophils and suggest that it may be a potential cellular phospholipid with CoA-IT inhibitor.

AM 15:49:54 AP 0

has been proposed to mediate the movement of arachidonate between phospholipid subclasses and influence the formation of arachidonic acid, a precursor of platelet-activating factor. To substantiate this hypothesis, the authors have developed two structurally distinct inhibitors, SK&F 98625 [diethyl 2-[3,4,5-trimethylimidazole-1-yl]heptane phosphonate] and SK&F 45905 [(trifluoromethyl)phenyl]ureido]-2-[(trifluoromethyl)phenyl]ureido]-4,5-dichlorobenzene sulfonic acid]. These compounds inhibit the capacity to block microsomal CoA-IT activity, the transacylation of 1-alkyl-2-linoleoyl-sn-glycerol-3-phosphate transfer of [¹⁴C]arachidonate from 1-acyl-lysophosphatidylcholine (lyso-PE). Both SK&F 98625 and SK&F 45905 (10⁻⁶-10⁻¹⁹ M) in these two assays. In contrast, these compounds had little or no effect on other lipid-modifying enzymes. CoA-dependent acyltransferase or acetyltransferase revealed that both SK&F 98625 and SK&F 45905 inhibit the enzyme and prevented the acylation of lysophosphatidylcholine in a competitive manner. In intact human neutrophils, both compounds completely blocked the movement of [¹⁴C]arachidonate from 1-alkyl-2-linoleoyl-sn-glycerol-3-phosphate (GPE). In contrast, these compounds did not alter the incorporation of free arachidonate into phospholipids in intact cells. This is the first report to demonstrate the importance of CoA-IT in arachidonate metabolism. These results provide further evidence that CoA-IT is a key enzyme in the large pool of -etl-lysophospholipids in human neutrophils and suggest that it may be a potential cellular phospholipid with CoA-IT inhibitor.

D' 342:259557

T.	Inhibitors of Co/ Independent	As. Glas	chl	movement of
	arthrid net ntr	leather ind	pho thol	man neutrophils
Al	Ch. lton, F. B. H.	Fonteh, A. N.	Sun	Hickey, Deirdre M.
	E. Torpne, T. e. J.	Haye	uth	Lisa A.; Heravi, Javid
	Winnler, A. J.			

CS Section on Pulmo. by and Cris... Carol Me... ...man Gray School of
Medicine, Winston-Salem, NC 7-704, 1

SO Biochemistry (1968), 34(16), 1541-1546
 EDEN: BIGHAM; I.S.I. 0006-2950

D Journal

LA English

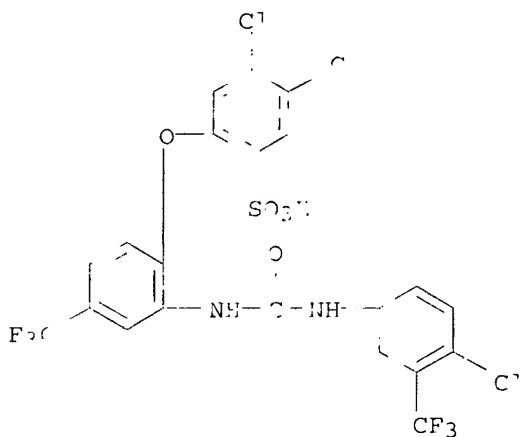
IT 162793-63-7

FL: BAC Biology activity of Effector, (not reverse); BIOL
Biology study;

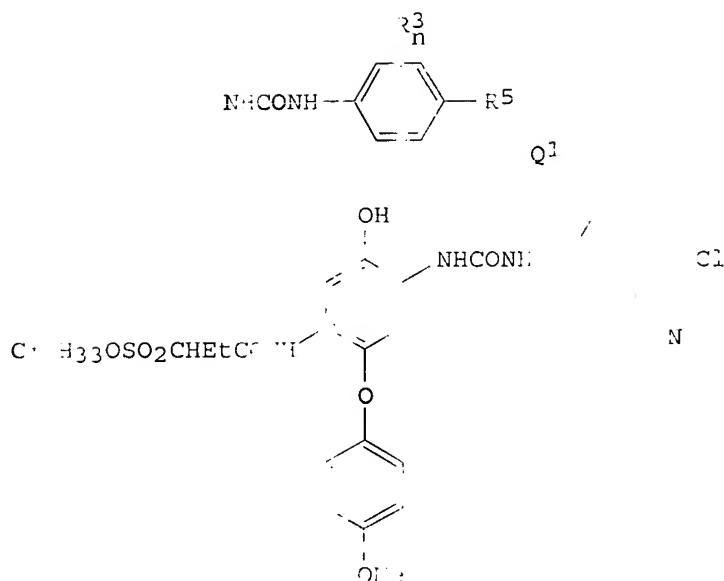
(inhibitor, a CoA-independent transacetylase, movement of
arachidonate to 1-ether-linked phospholipids in human neutrophils)

RN 162793-6. CAP

CN 4-azobenzoylformic acid, 4,5-dichloro-2-[4-(2,4,6-trifluoromethyl)phenyl]amino]-5-oxo-5H-pyridine-3-carboxylic acid
 (CI) (CA INDEX NAME)



L ANSWER 22 OF 22 C. PLUS COPYRT 1992 A
G



AE The title material contains a phenol cyan with a ureido group Q1 and 5-substituted R1 = (cyclo)alkyl, aryl, heterocycle; R2 = n = 1-4; R4 = H, alkyl, aryl, heterocycle. Thus, a soln. of the title cyan coupler in a red-sensitive AgBr emulsion then coated on photog. film, which gave fog-free printed

plate which is 2-substituted R1O O2R2CONH [Q2 = NR4, O; aryl; R3 = H, substituent; = 1 substituent except CN]. i.e. phthalate and EtOAc was mixed with a polymer ester support to give a high coloring property.

AP 1991:618758 CAPLUS
DI 115:218758
TI Silver halide color photographic emulsion ureido-substituted phenol cyan coupler
IN Nakayama, Noritaka; Masukawa, Toshiaki
P Konica Co., Japan
S Jpn. Kokai Tokkyo Koho, 11 pp.
CODEN: JKYYAF
DT Patent
LA Japanese
FAM.CNF 1

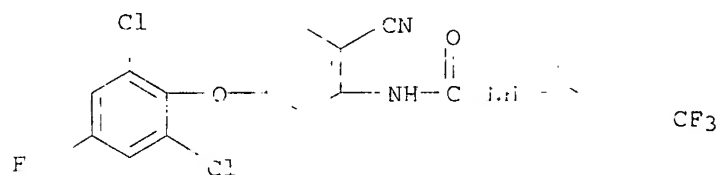
erica containing

PATENT NO.	KIND	DATE	AP	IOF	JO.	DATE
JP 0308024	A2	19910401	JP	-21	0	19890824

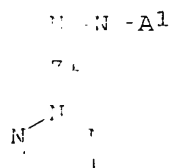
IT 136925-86-5
RL: USES (Uses)
(cyan coupler, for silver halide photographic film)
RM 136925-86-5 CAPLUS
CI Euranamide, 2-[(decylamino)sulfamoyl]-[1-[[[3-(trifluoromethyl)phenyl]amino]carbonyl]phenyl]- (9CI) (CA INDEX NAME)

ultra violet, prevention of fog
oxo-(4-methoxyphenoxy)-4-methylphenyl]- (9CI) (CA

R: BE, DE, FR, GB, IT, NL
 JP 02233650 A2 19890917
 PP: DE 1989-39028 19890106
 OS: MARPAT 11 367
 I: 132147-05-8
 RL: AGR (Agricultural use); B (biological activity); E (effect, except adverse); SPN (Synthetic preparation); B (biological study); PREP (Preparation); USES (Use); (prepn. of. as herbicide and as a growth regulator)
 R: 132147-05-8 CAPLUS
 C: Urea, N-[2-cyano-5-[2,6-dichloro-4-(trifluoromethyl)phenoxy]phenyl]- (9CI) IND (index)



L: ANSWER 24 OF 2 CAPLUS COPY: 1 002
 G



A: N=N-Z3 Z2-N=N-1

AB In the title photoreceptor having a photoconductive layer, the photoconductive pigment I (Z1, Z2, Z3 = Arylene, heterocyclic; A = methylene; A1, A2, A3 = coupler residue having phenolic OH)

A 1989:163564 CAPLUS

DI 110:163564

T. Electrophotographic photoreceptor containing pigments charge-generating trisazo

MI Miyazaki, Hajime; Takai, Hideki; Matsui, Yoichi

PA Canon K. K., Japan

SO Jpn. Kokai Tokkyo Koho, 27 pp.

CODEN: JEXXAF

DI Patent

L: Japanese

F: CNT 1

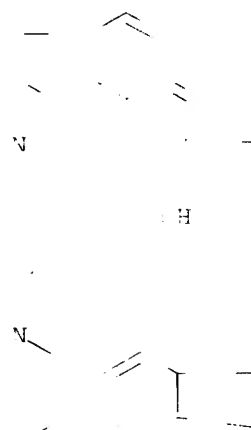
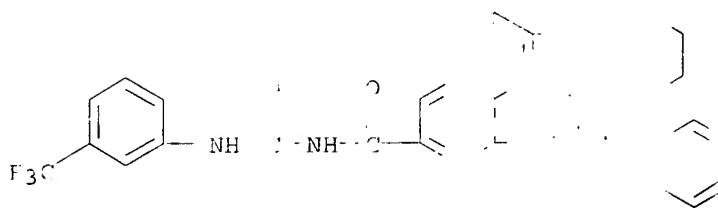
PATENT NO.	KIND	DATE	AP.	IO.	O.	DATE
JP 6328271	A2	19881114	JP	11	8	370515
JP 256080		19861201				
119956-25-8						
RL: USES (Uses)						

(electrophotog. charge-generating pig-
 119956-85-5 "APL" 1,1
 11H-Benzo[a]carbazole-3-carbonyl 1,1
 [[[[[3-(trifluoromethyl)phenyl]carbonyl
 benzo[a]carbazol-1-yl]amino]phenyl]car-
 4,1-phenylenebis[2-phenoxy-3-
 onyl]-C CA INDEX 1,1,1

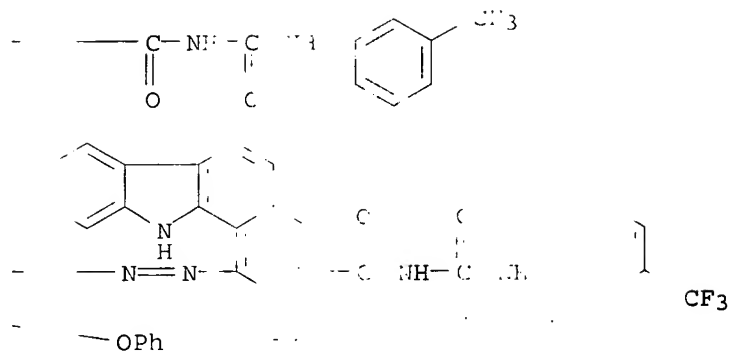
for improved sensitivity)

5-hydroxy-2-[4-[[2-hydroxy-3-
 amino]carbonyl]-11H-
 ne-5-dimethylbis[(2-phenoxy-
 4,1-phenylene]amino]carb

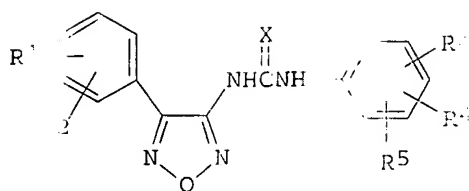
PAGE 1-A



PAGE 1-B



L ANSWER 20 00 2 PLUS 00 00 002
 G



I

A The title compds. [I; R1, R2 = H, halo, (halo)alkyl, (un)substituted aryl, optionally with 1 or 2 O, inter alia, or 2 O; R3, R4 = H, halo, (halo)alkyl, (halo)cycloalkyl; X = O, S] were prepared as pesticides and acaricides. 4-(2-chloro-5-(3,3,3-trifluoromethyl)-2-oxo-1-phenyl-1H-pyrazol-4-yl)-2-chloro-5-(3,3,3-trifluoromethyl)-2-oxo-1-phenyl-1H-pyrazole was hydrogenated over Raney Ni to give 4-(2-chloro-5-(3,3,3-trifluoromethyl)-2-oxo-1-phenyl-1H-pyrazol-4-yl)-2-chloro-5-(3,3,3-trifluoromethyl)-2-oxo-1-phenyl-1H-pyrazole. Tetranychus urticae eggs and larvae were killed by all stages, compared to 62% for the control.

A 1988:150483 CAT

D 108:150483

T Preparation of 1-phenyl-4-(4-chloro-2,3,3-trifluoromethyl-2-oxo-1-phenyl-1H-pyrazol-4-yl)-2-chloro-5-(3,3,3-trifluoromethyl)-2-oxo-1-phenyl-1H-pyrazole.

I Sirrenberg, Wilhelm; Mayold, Robert; Bayer A.-G., Fed. Rep. Ger.

P Bayer A.-G., Fed. Rep. Ger.

SC Ger. Offen., 29 pp.

CODEN: CNYXEX

D Patent

I German

F CNT 1

	PATENT NO.	IND	DATE	AP	MON	Q.	DATE
P	DE 3621662	A1	19880101	DE	67	62	19870708
	US 4853397	A1	19900101	US	67	62	19870625
	EP 253175	A2	19880101	EP	67	62	19870626
	EP 253175	A3	19900101				
	EP 253175	A1	19920101				
	R: CH3, CF3, C(CH3)3, DE, R, C(CH3)3, I, I						
	AT 4031	A1	19900101	AT	10	9	19870626
	JP 6301871	A1	19880101	JP	6		19870706
	DK 8703503	A1	19880101	DK	250		19870707
	HU 4417	A1	19880101	HU	306		19870707
	ZA 870118	A1	19880101	ZA	393		19870707

P DE 1984-3621662 19860101 19870708

C CASREAC 108:150483; MFI PAT 108:150483

I 113664-71-4P

RL: AG (agricultural use); L (logically adverse); SPH (Synthetic preparation); R (preparation); US (Use)

(preparation of as insecticide or acaricide)

R 113664-71-4 CAT

C 113664-71-4 CAT
UN 1, 1' [4-(2-chloro-5-(3,3,3-trifluoromethyl)-2-oxo-1-phenyl-1H-pyrazol-4-yl)-2-chloro-5-(3,3,3-trifluoromethyl)-2-oxo-1-phenyl-1H-pyrazole]
N 113664-71-4 CAT
CA INDEX NAME

alk (halo)alkoxy,
= substituted alkylene,
the Ph group with 1 or
y: (unsubstituted
and animal pests, esp.
clobutyl)nitrobenzene
iline was condensed
= H, R5 =
Tested against
II gave 100% kill of
anil insecticides.

as pesticides

as about

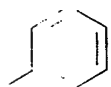
ativ y or effector, except
biological study; PREP

-3 (trifluoromethyl)phenyl]-
2 (oxadiazol-3-yl)- (9CI)

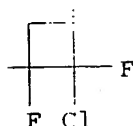
۴۰



3



I



1. addition of a polyether
 2. diisocyanate in
 3. to which monensin
 4. (fluoromethyl)-4,4'-
 5. use no. of combinations
 6. of monensin and
 7. of efficacy to the
 8. of
 9. with
 10.

L 3 1

Notion and

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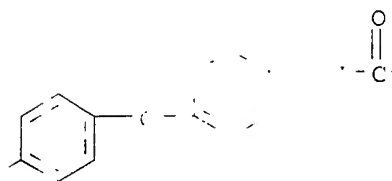
70

11

•

Point selected from C. The s. of

C urea, H-¹³C, flu [4] (4 p-methoxy)phenyl]-
 (DCI) (A) E)



L	ANSWER 77 OF 30	LUS	77 OF 30
A	Anilino RZC6H5 SO2) were present carboxamides, a chloroformate	R = C co nzer acy	77 OF 30 77 OF 30 77 OF 30 77 OF 30

lor pirazinyl, Z = O,
esp ureas, carbamates,
men isocyanates,

P 1984:5 810 CA
D 101:100 40
7 Synthes of pol al : : :
substituted and

an ... c des from

A Kempter, Gerhard verk
Ch. Sehl. Chem./33 aed sch "F
Foto:Ger. Sanssouci DR- De
S. Wiss. f. Paedag. Soch k
101-20
CODEN: JPHLAC 01

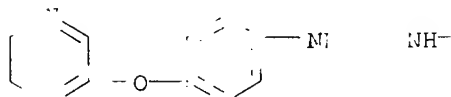
ebk' 1171 "

D Journal
L German
C CA 91645-5 101:11
I 91645-5 -5P
FL: SPH Synthet (prep. of) (P

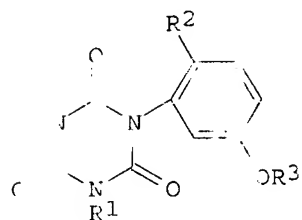
Pot 193, 27(1),

P 0160-77-5 CA
C Urea, N-[4-(3-methylbutyl)-2-pyridyl]-N'-methyl-N,N'-bis(2-oxoethyl)-
(CA) EX NAME

luo' me' ... r'henyl] - (9CI)



1 ANSWER 28 OF 12 PLUS 1 2 1
C



A Triazinetriones (R = substituted a
= H, alkyl, acyl, alkyl, morpho
substituted Ph] were prepared by the
P4CONCO (R4 = hal, alkyl, aryl).
(trifluoromethyl)phenoxyl, phenyl
ClCONCO to give 8. I = H, P
are effective herbicides. 3.0 kg

A 1083:408238 CAS
D 99:88238
T 1,3,5-Triazinone and th... for cont
I Parg, Adclif; Ham...; ...
P BASF A.-G., Fed. Rep.
S Ger. Offen., 55
CODEN: GWXXBX
D Patent
L German
F CIT 2

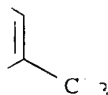
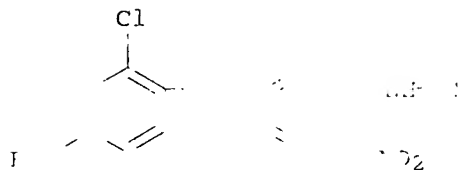
alkyl, cycloalkyl, Ph; R1
hal, cyano, NO2; R3 =
a phenylurea with
-[3-chloro-4-
thyl] was treated with
2, F = 2.4 Cl (F3C)C6H3). I

g un desired plant growth
o

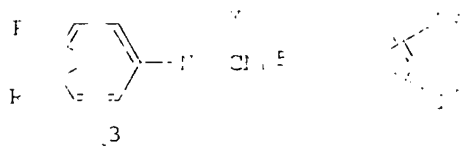
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P	DE 3147879	15	DE	3147	79 19811203
	EP 81110	5	EP	1108	9 19811124
	EP 81111	1			
	EP 81112	25			
	R: AT, BE, DE, FR, GB, IT, NL, PT, SE, CH, JP				
	JP 58103374	30		204	19811124
	CA 1135274	3	CA	4162	19811124
	AT 20528	1	AT	1108	9 19811124
	FR 3206046	1	FR	6946	19821130
	ZA 320607	1	ZA	885	19821202
	HU 30900	18	HU	3882	19821202
	HU 16831	13			
	US 451107	3	US	4620	4 19810128
F	DE 1981-3147879	13			
	DE 1981-3201229	5			
	EP 1981-10859	14			
	US 1982-46064				
C	CASREACT 99:88238				
I	80810-51-2				
	FL: F01 Reactant				
	(cyclocondensation of ... and ...)				
F	80810-51-2 CAS				
C	U.S. Pat. 3-10859				
	trifluoromethyl ...				

Int select * from ... 7/20

L: RCT 11/25/81
 (C) 11/25/81
 P 6607-16
 C 11/25/81
 (CA 11/25/81)



1 ANSWER OF 20 JULY 1964
C



Anticoccidial compns. for chicken or turkey control	feedstuff combinations	mixes for poultry such as
carbanilide I (R1, R2, and	H, halogen	polymer antibiotic and a
alkanoylamino, and alkyl	substituted	, NH, NO2, C1-6 alkyl, C2-4
C1-4 alkyl, R3, and R4	halogen	noxy, etc.; R4 and R5 = H or
etc.). Also, the mix of	amino	H2, C2-4 haloalkenyloxy,
trifluoromethyl carbamate	5393-2	o-3 and p-tro-5-
at 50 ppm effectively controls	coccidiosis	monosir [17090-79-8] each
infects with S. Typhimurium	and P. ne	broiler chicks

F 981:71 2 C.

I 34:71426

Anticoccolidial - posit cr

Callendo, Matt ; Emerson;

Plunkett, Clinton Albert

F Billy, 111 and 112, USA

S	Sur.	Pos.	or	pp.
1	1	1	1	1
2	2	2	2	2
3	3	3	3	3
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87	87	87	87	87

CDEN: XLD:

Patent

L. English

F .CMT 1

PATENT NO. _____ AND NO. _____ FILED ON _____ DATE _____

EP 15117 42 1 EP 4000 7 1000211

EE 15110	A3	15
EE 15111	A3	19

RE, PE, FF, C, U, NO,
S 421, 8, 19, U, 121, 790214

GB 104	99	13	GE	447	200211
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U 805 2 19 2 3 5546 00212

PH 5316 2 1

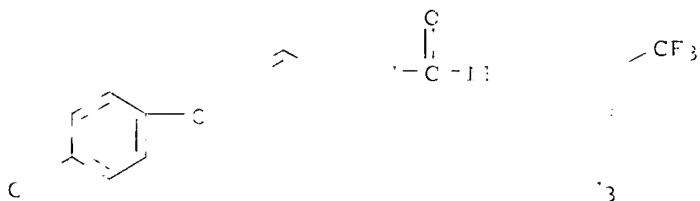
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HU 1850	1	19			
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FI 7148	1	19			
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US 40	1	19	US	100	820917
US 1975	1	19			
2063-6					

PL: BIO (bio) study
(anticoagulant) compn.
polyeth
phenyl - (chloromethoxy)phenyl]-



ANSWER: 30
For diazonium(s) prep
Highly light (t
R1 = e, Me, MeS, 4
+ [4-(3-CF3SCF3)
-ClCH2] or 4
sed in hot
field by react
solvents conta
1975:13900
82:1390
Highly light
Bettler, 1974
Bettler
Parbuer, 1968
Ger. 24
CPEN: 0000
alent
erman
C.T.
PATENT
ND
OP
E

1975:13900
82:1390
Highly light
Bettler, 1974
Bettler
Parbuer, 1968
Ger. 24
CPEN: 0000
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07/20/70

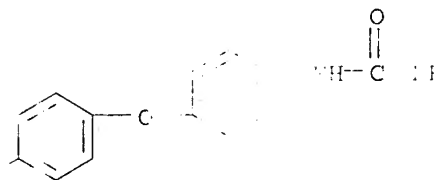
DE 2334355
2063-69-6P

DE 2334355 30706

RL: PREP (Preparation)
(manuf. of Helix)

2063-69-6-4-1
Urea, N-3,4-bis(4-fluorophenyl)-
(9CI) (CA 1971:1151)

phenyl 4-(4-fluorophenoxy)phenyl-



CF

ANSWER OF THE QUESTION
I, II, III, and IV are prepared by the method of Arion, Vimax, and Glabrat, and the method of 32.2 g. of 4-fluorophenylamine and 30 ml. of dioxane, give 4 g. of R3 = CF3, R4 = H, R5 = H, effective against Helix asna. "Twelve" 5 g. of Alternative method: 1-naphthylamine, 1.0 g. water-soluble urea, 0.5 g. compds. prepared by the method of Arion.

TEST 20

Tested by Helix, Arion, Vimax, Glabrat, and 1.3 g. of 4-fluorophenylamine and 30 ml. of dioxane, give 4 g. of R3 = CF3, R4 = H, R5 = H, effective against Helix asna. "Twelve" 5 g. of Alternative method: 1-naphthylamine, 1.0 g. water-soluble urea, 0.5 g. compds. prepared by the method of Arion.

Helix, Arion, Vimax, Glabrat, and 1.3 g. of 4-fluorophenylamine and 30 ml. of dioxane, give 4 g. of R3 = CF3, R4 = H, R5 = H, effective against Helix asna. "Twelve" 5 g. of Alternative method: 1-naphthylamine, 1.0 g. water-soluble urea, 0.5 g. compds. prepared by the method of Arion.

2069:49

71:9107

Urea and 4-fluorophenylamine

useful for

Helix asna

CIBA Ltd.

FR 1,910,710

CODEN: TRXMA

Patent

French

CNT 1

PATENT NO.

ID NO.

AP

CV

TE

FR 1,910,710

FR 1,910,710

20751-88-4

RL: SFM (Sulfonamide)
(prepn. of)

PREP

PREP

20751-88-4-1

Carbanilide, 3,4-bis(4-fluorophenyl)-

3,5-bis

3,5-bis

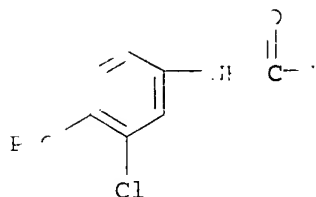
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INDEX

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Page 1

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FILE 'RLG STR' STEFF 155:43 ON 1

1 STP RE U C

1 13 S

1 365 S

FILE 'USPA FU' AT 15:5

1 17 S

FILE 'C PLUS' 13 ON 1

1 32 S

= s 15 and cancer

163322 CAN

22750 AM

170005 CAN

1 6 15 17

= d abs bib hits

1 ANSWER 1 OF 6

7 Chem. structure

kinase and in

pharmaceut

disorders

glycolytic

all pathwa

of oxygen

cancer and Al

1-phenylethy

1001:860012

1001:700

allosteric in

Abraham, L

James C. L

USA

U.S. Pat. 4,000

CODEN: 1770

1001:860012

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1001:860012

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P I 'S 1998-16113 99
1 89060-07-7

90324

PL: BSU (Biochem) ; HU (Therap) ; (pyruvate kinase)

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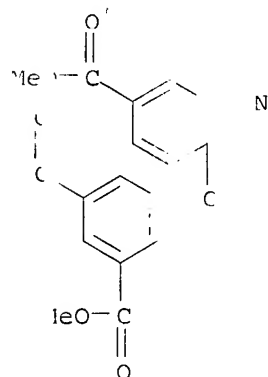
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        console.log('Average: ' + average);
    } else {
        console.log('No average calculated');
    }
}

// Example usage
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calculateAverage([1, 2, 3, 4, 5, NaN]); // Output: No average calculated
calculateAverage([1, 2, 3, 4, 5, NaN, NaN]); // Output: No average calculated

```

P 89060-C7-1 CA
C 1,3-Benzenedicarbonyl chloride (Mitsunobu)
(trifluoromethyl) phosphine oxide
(9CI) (A) N/A M

4-methyl-2-pentyl-13-bromopentadecanoic acid methyl ester



1 ANSWER 7 OF 6

C.

* PICTURE DIAGRAM OF ... AY ... LINE PRINT *

Title compd. 2,6-(CH₃)₂, 2-CH₃(CH₂)₅NHCO, 2-CH₃(CH₂)₆NHCO, m = 0-8; n = 0, and a therapeutic aid or prevent arteriosclerosis, cancers, as prepd.

7 0000:742094

I 33:296435

preparation of the product is useful in the treatment of hyperlipemia,

I arteriosclerosis of the coronary arteries; Fujita, Takeshi; 1940; 1941; 1942; 1943; 1944; 1945; 1946; 1947; 1948; 1949; 1950; 1951; 1952; 1953; 1954; 1955; 1956; 1957; 1958; 1959; 1960; 1961; 1962; 1963; 1964; 1965; 1966; 1967; 1968; 1969; 1970; 1971; 1972; 1973; 1974; 1975; 1976; 1977; 1978; 1979; 1980; 1981; 1982; 1983; 1984; 1985; 1986; 1987; 1988; 1989; 1990; 1991; 1992; 1993; 1994; 1995; 1996; 1997; 1998; 1999; 2000; 2001; 2002; 2003; 2004; 2005; 2006; 2007; 2008; 2009; 2010; 2011; 2012; 2013; 2014; 2015; 2016; 2017; 2018; 2019; 2020; 2021; 2022; 2023; 2024; 2025; 2026; 2027; 2028; 2029; 2030; 2031; 2032; 2033; 2034; 2035; 2036; 2037; 2038; 2039; 2040; 2041; 2042; 2043; 2044; 2045; 2046; 2047; 2048; 2049; 2050; 2051; 2052; 2053; 2054; 2055; 2056; 2057; 2058; 2059; 2060; 2061; 2062; 2063; 2064; 2065; 2066; 2067; 2068; 2069; 2070; 2071; 2072; 2073; 2074; 2075; 2076; 2077; 2078; 2079; 2080; 2081; 2082; 2083; 2084; 2085; 2086; 2087; 2088; 2089; 2090; 2091; 2092; 2093; 2094; 2095; 2096; 2097; 2098; 2099; 2100; 2101; 2102; 2103; 2104; 2105; 2106; 2107; 2108; 2109; 2110; 2111; 2112; 2113; 2114; 2115; 2116; 2117; 2118; 2119; 2120; 2121; 2122; 2123; 2124; 2125; 2126; 2127; 2128; 2129; 2130; 2131; 2132; 2133; 2134; 2135; 2136; 2137; 2138; 2139; 2140; 2141; 2142; 2143; 2144; 2145; 2146; 2147; 2148; 2149; 2150; 2151; 2152; 2153; 2154; 2155; 2156; 2157; 2158; 2159; 2160; 2161; 2162; 2163; 2164; 2165; 2166; 2167; 2168; 2169; 2170; 2171; 2172; 2173; 2174; 2175; 2176; 2177; 2178; 2179; 2180; 2181; 2182; 2183; 2184; 2185; 2186; 2187; 2188; 2189; 2190; 2191; 2192; 2193; 2194; 2195; 2196; 2197; 2198; 2199; 2200; 2201; 2202; 2203; 2204; 2205; 2206; 2207; 2208; 2209; 2210; 2211; 2212; 2213; 2214; 2215; 2216; 2217; 2218; 2219; 2220; 2221; 2222; 2223; 2224; 2225; 2226; 2227; 2228; 2229; 2230; 2231; 2232; 2233; 2234; 2235; 2236; 2237; 2238; 2239; 2240; 2241; 2242; 2243; 2244; 2245; 2246; 2247; 2248; 2249; 2250; 2251; 2252; 2253; 2254; 2255; 2256; 2257; 2258; 2259; 2260; 2261; 2262; 2263; 2264; 2265; 2266; 2267; 2268; 2269; 2270; 2271; 2272; 2273; 2274; 2275; 2276; 2277; 2278; 2279; 2280; 2281; 2282; 2283; 2284; 2285; 2286; 2287; 2288; 2289; 2290; 2291; 2292; 2293; 2294; 2295; 2296; 2297; 2298; 2299; 2300; 2301; 2302; 2303; 2304; 2305; 2306; 2307; 2308; 2309; 2310; 2311; 2312; 2313; 2314; 2315; 2316; 2317; 2318; 2319; 2320; 2321; 2322; 2323; 2324; 2325; 2326; 2327; 2328; 2329; 2330; 2331; 2332; 2333; 2334; 2335; 2336; 2337; 2338; 2339; 2340; 2341; 2342; 2343; 2344; 2345; 2346; 2347; 2348; 2349; 2350; 2351; 2352; 2353; 2354; 2355; 2356; 2357; 2358; 2359; 2360; 2361; 2362; 2363; 2364; 2365; 2366; 2367; 2368; 2369; 2370; 2371; 2372; 2373; 2374; 2375; 2376; 2377; 2378; 2379; 2380; 2381; 2382; 2383; 2384; 2385; 2386; 2387; 2388; 2389; 2390; 2391; 2392; 2393; 2394; 2395; 2396; 2397; 2398; 2399; 2400; 2401; 2402; 2403; 2404; 2405; 2406; 2407; 2408; 2409; 2410; 2411; 2412; 2413; 2414; 2415; 2416; 2417; 2418; 2419; 2420; 2421; 2422; 2423; 2424; 2425; 2426; 2427; 2428; 2429; 2430; 2431; 2432; 2433; 2434; 2435; 2436; 2437; 2438; 2439; 2440; 2441; 2442; 2443; 2444; 2445; 2446; 2447; 2448; 2449; 2450; 2451; 2452; 2453; 2454; 2455; 2456; 2457; 2458; 2459; 2460; 2461; 2462; 2463; 2464; 2465; 2466; 2467; 2468; 2469; 2470; 2471; 2472; 2473; 2474; 2475; 2476; 2477; 2478; 2479; 2480; 2481; 2482; 2483; 2484; 2485; 2486; 2487; 2488; 2489; 2490; 2491; 2492; 2493; 2494; 2495; 2496; 2497; 2498; 2499; 2500; 2501; 2502; 2503; 2504; 2505; 2506; 2507; 2508; 2509; 2510; 2511; 2512; 2513; 2514; 2515; 2516; 2517; 2518; 2519; 2520; 2521; 2522; 2523; 2524; 2525; 2526; 2527; 2528; 2529; 2530; 2531; 2532; 2533; 2534; 2535; 2536; 2537; 2538; 2539; 2540; 2541; 2542; 2543; 2544; 2545; 2546; 2547; 2548; 2549; 2550; 2551; 2552; 2553; 2554; 2555; 2556; 2557; 2558; 2559; 2560; 2561; 2562; 2563; 2564; 2565; 2566; 2567; 2568; 2569; 2570; 2571; 2572; 2573; 2574; 2575; 2576; 2577; 2578; 2579; 2580; 2581; 2582; 2583; 2584; 2585; 2586; 2587; 2588; 2589; 2590; 2591; 2592; 2593; 2594; 2595; 2596; 2597; 2598; 2599; 2600; 2601; 2602; 2603; 2604; 2605; 2606; 2607; 2608; 2609; 2610; 2611; 2612; 2613; 2614; 2615; 2616; 2617; 2618; 2619;

F. Bankyo Company, Ltd.,

S. 100 CT Int. Appl. 20 11.

CODEN: FJXXD2

Patent

1 Japanese

P. CUT 1

PATENT NO. 2,145,147

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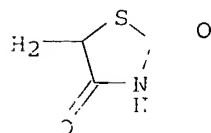
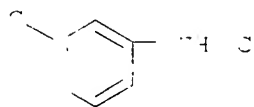
Print selected for [redacted] 10/15/93

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Print selected from

References 1/17/2012

W: AU, BR	CA, CH	DE, FR	NO	PL, RU, TR,
US, JP				
RW: AT, BE	CH, DE	FR, IT,		LU, MC, NL,
T, S				
JP 2000-1167	2000	10		406
JP 1167	2000	10		406
R: AU, BR		S, FR,		SE, MC, PT,
BR 20000000004	2000	04	0501	406
NO 2001001007	2001			11005
F JP 1999-0001	1999	01		
WO 2000-100016	2000	01		
C MARPAT 103:000				
I 301548-72-2P				
EL: BAC (biological	study, in vitro	effect		BSU (Biological
study, in vitro	study, in vitro	study, in vitro		therapeutic use);
BIOL (Biological	study, in vitro	study, in vitro		
(preparation)	study, in vitro	study, in vitro		hyperlipemia,
artefact	study, in vitro	study, in vitro		
F 301548-72-2C				
C Urea, N-[3-(2-	thio-1,3,4-	thio-1,3,4-		noxy)methyl]-1-
methyl-1H-benz-	thio-1,3,4-	thio-1,3,4-		thyl)phenyl]-
(2CI) (A) (N)	thio-1,3,4-	thio-1,3,4-		

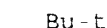


B

F CNT 16	THIOPHENE	THIOPHENE	THIOPHENE	FLCORN
	THIOPHENE	THIOPHENE	THIOPHENE	
I ANSWER 3	THIOPHENE	THIOPHENE	GHT 100	
C				

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References 17:01 Page



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esp. Ph or
first one
q = 1-3; B =
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ford the urea

FIGURE 1. COT 1

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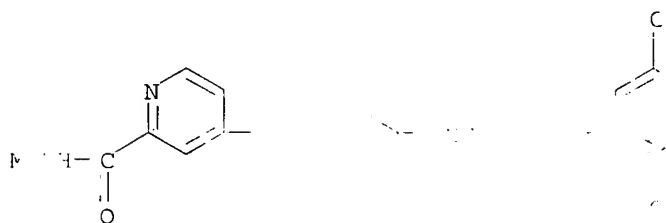
10202

77 47' 15" 15

MS 1999 .2.2

Patent selected from WO 99/0012

WO 2000 50000
C MARPAT
284461-4
RL: BAC (Biological)
study, unclassified; THU
Therapeutic (Biological)
Reactant (Biological)
(prepn. of)
kinase (Biological)
F 284461-4-3
C 2-Pyridylamino
carbonyl (Biological)



12
ANSWER 4 ON 6
A method of treating a
comprises a
2-thienyl; B
arom. substituted
tetrahydro
tirred
tetrahydro
inhibitor
A 1999:427
E 131:586
T Preparation of
D Miller, Scott
Timothy Bruce
Bill E.; Gunn
Robert; Wood
F Bayer Corporation
S CT Int. Ag.
CODEN: 131:586
E Patent
I English
F 1
PATENT 12
F WO 9932467
W: AU, A
RW: CH, G
SZ, LU, M

under
pyridyl,
6-membered
butyl-2-(3-
ocanate were
16 compds.
Lowinger,
ger A.; Wood,
Sibley,

WO 9932467
W: AU, A
RW: CH, G
SZ, LU, M
CZ, DE,
IN, IS, JP,
MG, MK, MN,
TJ, TM,
RU, TJ, TM
DE, DK, ES,
CF, CG, CI,

Patent selected from WO 99/0012

It selected from

CA 2315774

AU 9919350

EP 1042305

R: A

JP 2001576276

US 1997-15774

WO 1998-13272

CARPAT 131-50

28399-63-1P

AB: BAC

study, and

BIOL (BIO)

(prepr)

28399-63-1P

Urea, N-methyl

methoxy

981 22

981 22

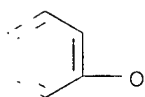
981 22

SE, MC, PT,

81 22

; B U (Biological
therapeutic use);

M



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RECORD

I

ANSWER 5

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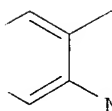
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heteroaryl]

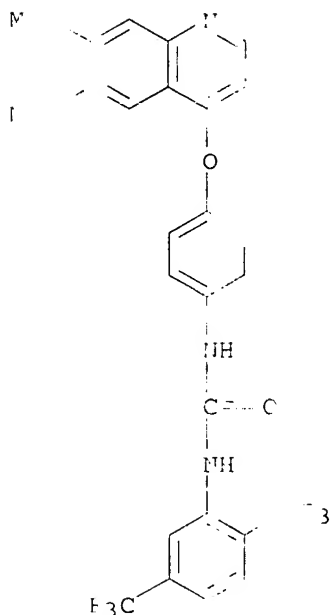
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ANSWER C OF ...
 ET-18-O-CH₃ ...
 antiproliferative ...
 both in vitro and ...
 mechanism ...
 transacylase (C ...
 specific phosph ...
 cells. ET-18-O ...
 and kinetic ana ...
 lyso-phospholip ...
 the connect ...
 using several st ...
 other inhibitory ...
 thymidine incor ...
 human HL-60 cell ...
 by ET-18-O-CH₃ ...
 necrosis factor ...
 necrosis factor ...
 model revealed ...
 inhibitors, ...
 apoptosis. ...
 participate ...
 phospholipase A ...
 demonstrated th ...
 proliferation ...
 996:70-144 (AF ...
 26:166-148 ...
 Inhibitors of ...
 human HL-60 cell ...
 Linker, Jan ...
 Mayer, Ruth ...
 Dep. Immunol ...
 of Prun ...
 Journal ...
 96, 279(2),

Print selected for [unclear] [unclear] [unclear]

156-966

MODEN: JPET.

Williams & [unclear]

Journal

English

162793-63-

RL: BAC (Biochemical)

study, unclassified

(inhibitor of

HL-60 cells)

162793-63-7

benzenesulfonate

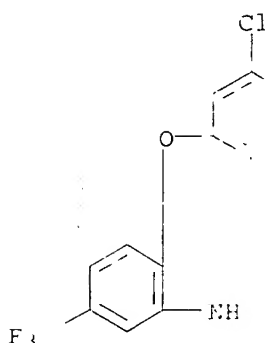
(trifluoromethyl)

(OCI) (C) E

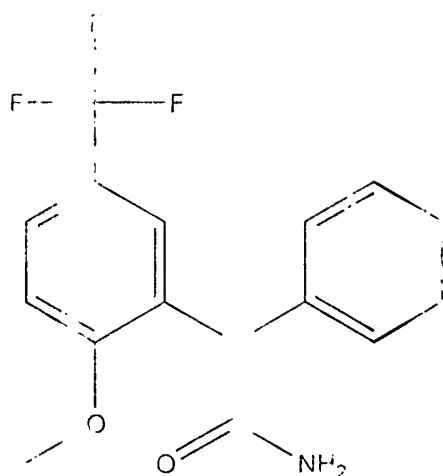
RSU (Biological

apoptosis in human

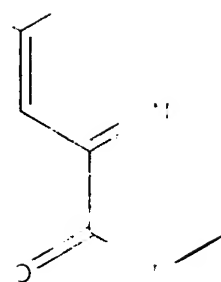
[4-methylphenoxy]-



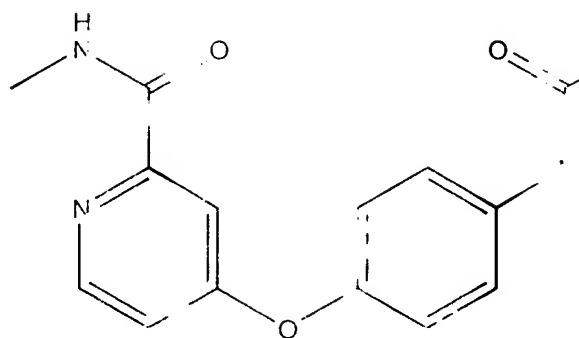
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N-(2-methoxy-5-(trifluoromethyl)phenyl)-N'-4-(2-methyl-5-pyridyloxy)phenylurea



N-(2-methoxy-5-(trifluoromethyl)phenyl)-N'-4-(2-methyl-5-pyridyloxy)phenylurea



N-(2-methoxy-5-(trifluoromethyl)phenyl)-N'-4-(2-(N-methyl-5-pyridyloxy)phenyl)urea

L4 ANSWER 11 OF 17 USPATFULL

AB This invention relates to the novel pharmaceutical compositions of Formulas (I) and (II) each of which comprises a compound of Formula (I) or (II) and a pharmaceutically acceptable diluent or carrier.

This invention also relates to a method of treating or reducing inflammation in a mammal in need thereof, which comprises administering to said mammal an effective amount of a compound or composition of Formula (I) or (II).

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AN 95:105872 USPATFULL

TI Anti-inflammatory compounds

IN Dixon, James S., Malvern, PA, United States
Hall, Ralph F., Villanova, PA, United States
Marshall, Lisa A., Wyndmoor, PA, United States
Chilton, III, Floyd H., Pilot Mountain, NC, United States
Mayer, Ruth J., Wayne, PA, United States

PA Winkler, James D., Fort Washington, PA, United States
SmithKline Beecham Corp., Philadelphia, PA, United States (U.S. corporation)

PI US 5470882 19951128

AI US 1994-252716 19940602 (8)

DT Utility

FS Granted

EXNAM Primary Examiner: Dees, Jose G.; Assistant Examiner: Conrad, III, Joseph M.

LREP Dinner, Dara L., Venetianer, Stephen, Lentz, Edward T.

CLMN Number of Claims: 5

ECL Exemplary Claim: 1

DRWN No Drawings

LN.CNT 1612

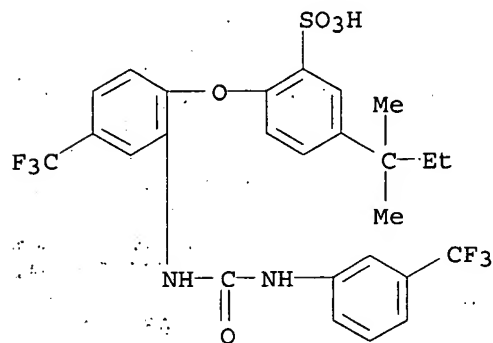
CAS INDEXING IS AVAILABLE FOR THIS PATENT.

IT 447-64-3

(anti-inflammatory benzenesulfonic acid derivs., their prepn., and their activity)

RN 447-64-3 USPATFULL

CN Benzenesulfonic acid, 5-(1,1-dimethylpropyl)-2-[4-(trifluoromethyl)-2-[[[3-(trifluoromethyl)phenyl]amino]carbonyl]amino]phenoxy]- (9CI) (CA INDEX NAME)



L4 ANSWER 12 OF 17 USPATFULL

AB This invention relates to the novel compounds and pharmaceutical compositions of Formula (I).

This invention also relates to a method of treating or reducing inflammation in a mammal in need thereof, which comprises administering to said mammal an effective amount of a compound or composition of Formula (I).

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AN 95:80325 USPATFULL

TI Anti-inflammatory compounds

IN Adams, Jerry L., Wayne, PA, United States
Hall, Ralph F., Villanova, PA, United States
Seibel, George L., Wayne, PA, United States

PA SmithKline Beecham Corp., Philadelphia, PA, United States (U.S. corporation)

PI US 5447957 19950905

AI US 1994-252851 19940602 (8)

DT Utility

FS Granted

EXNAM Primary Examiner: Dees, Jose G.; Assistant Examiner: Barts, Samuel

LREP Dinner, Dara L., Venetianer, Stephen, Lentz, Edward T.

CLMN Number of Claims: 12

ECL Exemplary Claim: 1

DRWN No Drawings

LN.CNT 1726

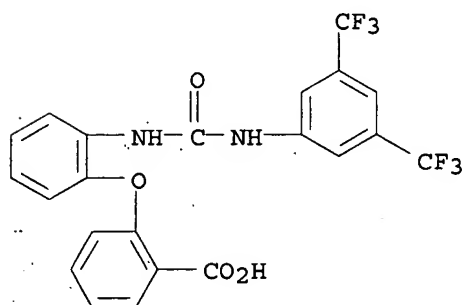
CAS INDEXING IS AVAILABLE FOR THIS PATENT.

IT 171103-10-9P

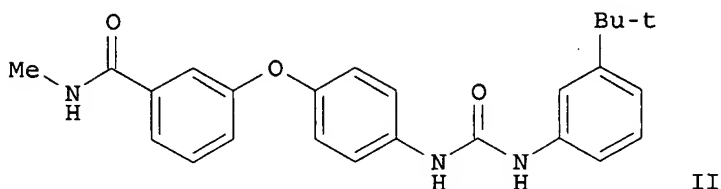
(antiinflammatory (ureidophenoxy)benzoic acids and derivs. as inhibitors of phospholipase A2 and CoA-independent transacylase)

RN 171103-10-9 USPATFULL

CN Benzoic acid, 2-[2-[[[3,5-bis(trifluoromethyl)phenyl]amino]carbonyl]amino]phenoxy]- (9CI) (CA INDEX NAME)



L5 ANSWER 7 OF 32 CAPLUS COPYRIGHT 2002 ACS
GI



AB This invention relates to the prepn. and use of (hetero)aryl ureas ANHCONHB [I; A = L(ML1)q; L = 5- or 6-membered (hetero)aryl, esp. Ph or pyridinyl; M = bridging group; L1 = (hetero)aryl with at least one (un)substituted sulfamoyl, carboxy, or carbamoyl substituent; q = 1-3; B = certain (un)substituted mono- to tricyclic aryl or heteroaryl groups] for the treatment of raf mediated diseases, such as cancer (no data). Approx. 100 invention compds. and numerous intermediates were prepd. For instance, 3-tert-butylaniline was coupled with bis(trichloromethyl)carbonate to form the isocyanate, followed by addn. of 4-(3-N-methylcarbamoylphenoxy)aniline (prepn. given) to afford the urea II.

AN 2000:493516 CAPLUS

DN 133:120157

TI Preparation of .omega.-carboxy(hetero)aryl substituted diphenyl ureas as raf kinase inhibitors

IN Riedl, Bernd; Dumas, Jacques; Khire, Uday; Lowinger, Timothy B.; Scott, William J.; Smith, Roger A.; Wood, Jill E.; Monahan, Mary-Katherine; Natero, Reina; Renick, Joel; Sibley, Robert N.

PA Bayer Corporation, USA

SO PCT Int. Appl., 120 pp.

CODEN: PIXXD2

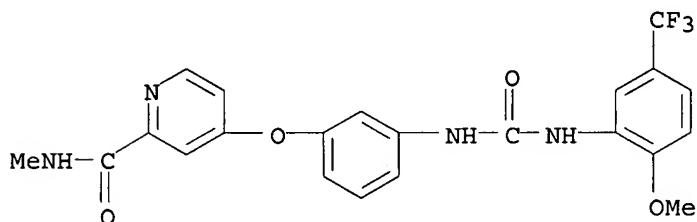
DT Patent

LA English

FAN.CNT 1

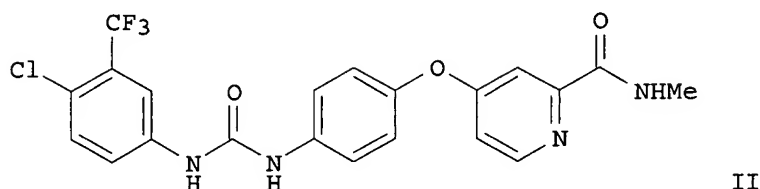
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PI	WO 2000042012	A1	20000720	WO 2000-US648	20000112
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	RW:	GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG			
	EP 1140840	A1	20011010	EP 2000-903239	20000112
	R:	AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO			
	US 2001011135	A1	20010802	US 2001-773659	20010202
	US 2001011136	A1	20010802	US 2001-773675	20010202
	US 2001016659	A1	20010823	US 2001-773672	20010202

	US 2001027202	A1	20011004	US 2001-773658	20010202
	US 2001034447	A1	20011025	US 2001-773604	20010202
	NO 2001003463	A	20010912	NO 2001-3463	20010712
	US 2002042517	A1	20020411	US 2001-948915	20010910
PRAI	US 1999-115877P	P	19990113		
	US 1999-257266	A2	19990225		
	US 1999-425228	A2	19991022		
	WO 2000-US648	W	20000112		
OS	MARPAT 133:120157				
IT	284461-42-3P				
	RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses)				
	(prepn. of .omega.-carboxy(hetero)aryl substituted di-Ph urea raf kinase inhibitors by reacting arylisocyanates with arylamines)				
RN	284461-42-3	CAPLUS			
CN	2-Pyridinecarboxamide, 4-[3-[[[2-methoxy-5-(trifluoromethyl)phenyl]amino]carbonyl]amino]phenoxy]-N-methyl- (9CI) (CA INDEX NAME)				



RE.CNT 12 THERE ARE 12 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 8 OF 32 CAPLUS COPYRIGHT 2002 ACS
GI



AB The title compds. ADB [I; D = NHCONH; A = substituted moiety of up to 40 carbon atoms of the formula L(ML1)q (wherein L = 5-6 membered cyclic structure; L1 = substituted cyclic moiety having at least 5 members; M = bridging group having at least one atom; q = 1-3; each of L and L1 contains 0-4 members of the group consisting of N, O and S); B = (un)substituted up to tricyclic aryl or heteroaryl moiety of up to 30 carbon atoms with at least one 6-member cyclic structure bound directly to D contg. 0-4 members of the group consisting of N, O and S], useful in treating p38 mediated diseases, were prepd. E.g., a multi-step synthesis of the urea II which showed IC50 of 1-10 .mu.M against p38, was given. Compds. I are effective at 0.01-200 mg/kg/day (oral administration).

AN 2000:493376 CAPLUS

DN 133:120155

TI Preparation of .omega.-carboxy aryl substituted diphenyl ureas as p38 kinase inhibitors

IN Riedl, Bernd; Dumas, Jacques; Khire, Uday; Lowinger, Timothy B.; Scott, William J.; Smith, Roger A.; Wood, Jill E.; Monahan, Mary-Katherine; Natero, Reina; Renick, Joel; Sibley, Robert N.

PA Bayer Corporation, USA

SO PCT Int. Appl., 148 pp.
CODEN: PIXXD2

DT Patent

LA English

FAN.CNT 2

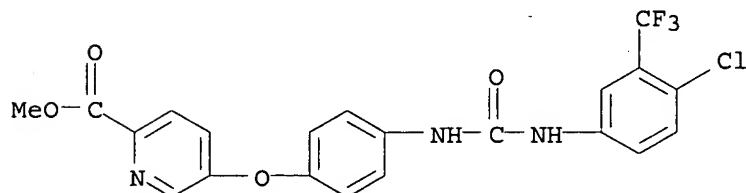
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PI	WO 2000041698	A1	20000720	WO 2000-US768	20000113
	W:				
	AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU, CZ, DE, DK, DM, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM				
	RW:				
	GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG				
	EP 1158985	A1	20011205	EP 2000-905597	20000113
	R:				
	AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO				
PRAI	US 1999-115878P	P	19990113		
	US 1999-257265	A2	19990225		
	US 1999-425229	A2	19991022		
	WO 2000-US768	W	20000113		
OS	MARPAT 133:120155				
IT	284461-86-5P				

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses)

(prepn. of .omega.-carboxy aryl substituted di-Ph ureas as p38 kinase inhibitors)

RN 284461-86-5 CAPLUS

CN 2-Pyridinecarboxylic acid, 5-[4-[[[4-chloro-3-(trifluoromethyl)phenyl]amino]carbonyl]amino]phenoxy]-, methyl ester (9CI)
(CA INDEX NAME)



RE.CNT 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 10 OF 32 CAPLUS COPYRIGHT 2002 ACS

AB A method of treating a p-38 mediated disease other than cancer comprises administration of BNHCONHA [A = (substituted) Ph, pyridyl, 2-thienyl; B = (substituted) aryl, heteroaryl contg. .gtoreq.1 6-membered arom. structure contg. 0-4 N, O, or S atoms]. Thus, 5-tert-butyl-2-(3-tetrahydrofuranyloxy)aniline (prepn. given) and p-tolyl isocyanate were stirred 8 h in PhMe to give 75% N-(5-tert-butyl-2-(3-tetrahydrofuranyloxy)phenyl)-N'-(4-methylphenyl)urea. Title compds. inhibited p38 kinase with IC50 = 1-10 .mu.M.

AN 1999:421667 CAPLUS

DN 131:58659

TI Preparation of diaryl ureas as inhibitors of p38 kinase.

IN Miller, Scott; Osterhout, Martin; Dumas, Jacques; Khire, Uday; Lowinger, Timothy Bruno; Riedl, Bernd; Scott, William J.; Smith, Roger A.; Wood, Jill E.; Gunn, David; Hatoum-Mokdad, Holia; Rodriguez, Mareli; Sibley, Robert; Wang, Ming

PA Bayer Corporation, USA

SO PCT Int. Appl., 107 pp.

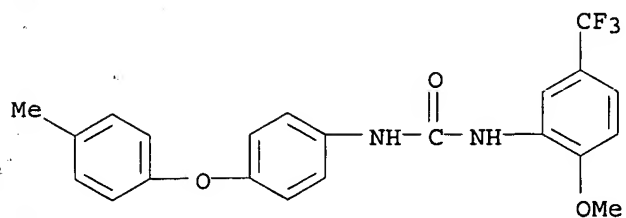
CODEN: PIXXD2

DT Patent

LA English

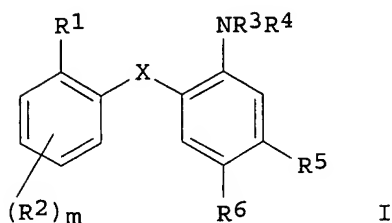
FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	WO 9932463	A1	19990701	WO 1998-US27265	19981222
	W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, UZ, VN, YU, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, RW: GH, GM, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG				
	CA 2315715	AA	19990701	CA 1998-2315715	19981222
	AU 9919399	A1	19990712	AU 1999-19399	19981222
	EP 1042305	A1	20001011	EP 1998-964221	19981222
	R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO				
	JP 2001526276	T2	20011218	JP 2000-525400	19981222
PRAI	US 1997-995749	A	19971222		
	WO 1998-US27265	W	19981222		
OS	MARPAT 131:58659				
IT	228399-63-1P				
	RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (prepn. of diaryl ureas as inhibitors of p38 kinase)				
RN	228399-63-1 CAPLUS				
CN	Urea, N-[2-methoxy-5-(trifluoromethyl)phenyl]-N'-[4-(4-methylphenoxy)phenyl]- (9CI) (CA INDEX NAME)				



RE.CNT 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 19 OF 32 CAPLUS COPYRIGHT 2002 ACS
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AB This invention relates to the novel compds. and pharmaceutical compns. of formula I wherein R1 is (CH2)nOH or (CH2)nCO2R8 ; n is 0 or an integer having a value of 1; X is oxygen or sulfur; R2 is hydrogen, halogen, optionally substituted C1-8 alkyl, or C1-8 alkoxy; m is an integer having a value of 1 or 2; R3 is C(O)R7 ; R4 is hydrogen, or C1-4 alkyl; R5 is hydrogen, halogen, CF3, CH3, (CH2)tCO2R9, or (CH2)tOH; t is 0 or an integer having a value of 1 or 2; R6 is hydrogen or halogen; R7 is NR9R10 ; R8 is hydrogen or C1-4 alkyl; R9 is hydrogen or C1-4 alkyl; R10 is hydrogen, optionally substituted aryl, optionally substituted arylC1-2 alkyl, optionally substituted C1-8 alkyl, or together R9 and R10 with the nitrogen to which they are attached form a 5 to 7 membered satd. or unsatd. ring which may optionally comprise an addnl. heteroatom selected from O/N or sulfur; or a pharmaceutically acceptable salt thereof. This invention also relates to a method of treating or reducing inflammation in a mammal in need thereof, which comprises administering to said mammal an effective amt. of a compd. or compn. of I. Thus, e.g., benzhydrol 2-[2-[3-(4-bromophenyl)ureido]-4-(trifluoromethyl)phenoxy]benzoate (prepn. given) was hydrogenated over 10% Pd/C to afford 2-[2-[3-(4-bromophenyl)ureido]-4-(trifluoromethyl)phenoxy]benzoic acid which inhibited PLA2 and CoA-IT at 50 .mu.M or less.

AN 1995:838690 CAPLUS

DN 124:8418

TI Antiinflammatory (ureidophenoxy)benzoic acids and derivatives as inhibitors of phospholipase A2 and CoA-independent transacylase

IN Adams, Jerry L.; Hall, Ralph F.; Seibel, George L.

PA SmithKline Beecham Corp., USA

SO U.S., 17 pp.

CODEN: USXXAM

DT Patent

LA English

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	US 5447957	A	19950905	US 1994-252851	19940602
	WO 9533460	A1	19951214	WO 1995-US6680	19950602
	W: JP, US				
	RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE				
PRAI	US 1994-252851		19940602		
OS	MARPAT 124:8418				
IT	171103-10-9P				

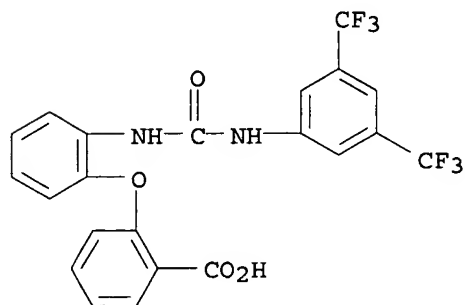
RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic

preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

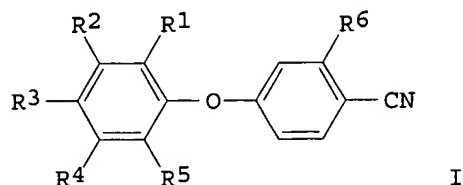
(antiinflammatory (ureidophenoxy)benzoic acids and derivs. as inhibitors of phospholipase A2 and CoA-independent transacylase)

RN 171103-10-9 CAPLUS

CN Benzoic acid, 2-[2-[[[3,5-bis(trifluoromethyl)phenyl]amino]carbonyl]amino]phenoxy]- (9CI) (CA INDEX NAME)



L5 ANSWER 23 OF 32 CAPLUS COPYRIGHT 2002 ACS
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AB The title compds. [I; R1 = H, cyano, CF3; R2, R4, R5 = H, halo; R3 = halo, CF3, CF3O, CF3SO2; R6 = NR7R8, CH2CHR11CO2R12; R7, R8 = H, alkoxy-carbonyl-ethyl, COR9, SO2R10; R9 = (un)substituted alkyl, alkenyl, alkynyl, Ph(CH2), naphthyl, pyridyl, furyl, PhS, alkylamino, etc.; R10 = (un)substituted alkyl, Ph, naphthyl, pyridyl, thienyl; R11 = H, halo; R12 = alkyl] were prep'd. as herbicides and plant growth regulators (no data), e.g., by etherification of amino(hydroxy)benzonitriles with halobenzenes. Thus, 3,4,5-trichlorobenzotrifluoride in DMSO was added dropwise to a pre-stirred mixt. of 2-amino-4-hydroxybenzonitrile and NaOH in DMSO and the whole was stirred for 5 h at 50.degree. and 2 h at 90.degree. to give 85% title compd. I (R1 = R5 = Cl, R2 = R4 = H, R3 = CF3, R6 = NH2).

AN 1991:101367 CAPLUS

DN 114:101367

TI Preparation of phenoxybenzonitriles as herbicides and plant growth regulators

IN Busse, Ulrich; Santel, Hans Joachim; Schmidt, Robert R.; Luerksen, Klaus; Strang, Harry

PA Bayer A.-G., Fed. Rep. Ger.

SO Eur. Pat. Appl., 31 pp.

CODEN: EPXXDW

DT Patent

LA German

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	EP 379915	A1	19900801	EP 1990-100701	19900113
	R: BE, CH, DE, FR, GB, IT, LI, NL				
	JP 02233655	A2	19900917	JP 1990-11973	19900123
PRAI	DE 1989-3902288		19890126		

OS MARPAT 114:101367

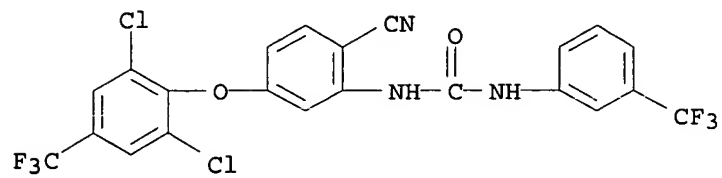
IT 132147-05-8P

RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses)

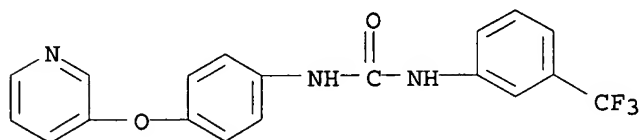
(prepn. of, as herbicide and plant growth regulator)

RN 132147-05-8 CAPLUS

CN Urea, N-[2-cyano-5-[2,6-dichloro-4-(trifluoromethyl)phenoxy]phenyl]-N'-[3-(trifluoromethyl)phenyl]- (9CI) (CA INDEX NAME)



L5 ANSWER 27 OF 32 CAPLUS COPYRIGHT 2002 ACS
AB Anilines RZC6H4NH2 (R = heteroaryl, e.g., 6-chloro-3-pyridazinyl, Z = O, SO₂) were prepd. and converted into their corresponding ureas, carbamates, carboxamides, and benzenesulfonamides by treatment with isocyanates, chloroformates, and acyl halides, resp.
AN 1984:510849 CAPLUS
DN 101:110849
TI Synthesis of potential plant protective agents and pesticides from substituted anilines
AU Kempter, Gerhard; Beerbalk, H. D.
CS Sekt. Chem./Biol., Paedagog. Hochsch. "Karl Liebknecht", Potsdam-Sanssouci, DDR-1500, Ger. Dem. Rep.
SO Wiss. Z. Paedagog. Hochsch. "Karl Liebknecht" Potsdam (1983), 27(1), 101-20
CODEN: WPKLAO; ISSN: 0138-290X
DT Journal
LA German
OS CASREACT 101:110849
IT 91619-55-5P
RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. of)
RN 91619-55-5 CAPLUS
CN Urea, N-[4-(3-pyridinyloxy)phenyl]-N'-[3-(trifluoromethyl)phenyl]- (9CI) (CA INDEX NAME)



L6 ANSWER 1 OF 6 CAPLUS COPYRIGHT 2002 ACS

AB Chem. structures have been identified which allosterically modify pyruvate kinase and inhibit enzymic activity. These compds. can be used as pharmaceuticals in the treatment of a wide variety of diseases and disorders where influencing metabolic processes is beneficial, e.g. the glycolytic pathway, all pathways which use ATP as an energy source, and all pathways which involve 2,3-diphosphoglycerate related to the delivery of oxygen by modifying Hb's oxygen affinity, treatments of tumor and cancer and Alzheimer's disease. Prepn. of e.g. 2-phenylethyloxy-5-formylbenzoic acid is described.

AN 2001:869018 CAPLUS

DN 136:700

TI Allosteric inhibitors of pyruvate kinase for therapeutic use

IN Abraham, Donald J.; Wang, Changging; Danso-Danquah, Richmond; Burnett, James C.; Joshi, Gajanan S.; Hoffman, Steven J.

PA USA

SO U.S. Pat. Appl. Publ., 15 pp., Cont.-in-part of U.S. 6,214,879.

CODEN: USXXCO

DT Patent

LA English

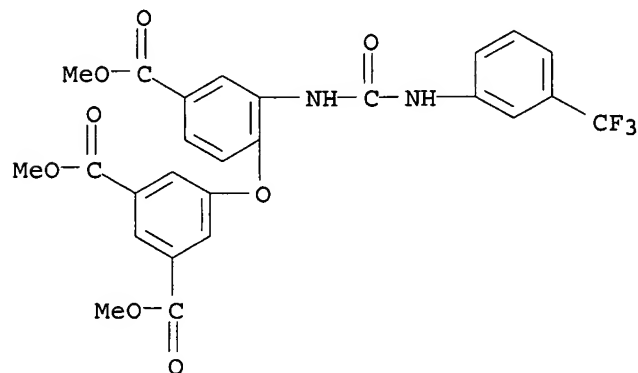
FAN.CNT 2

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	US 2001046997	A1	20011129	US 2001-799873	20010307
	US 6214879	B1	20010410	US 1998-46643	19980324
PRAI	US 1998-46643	A2	19980324		
IT	289060-07-7				

RL: BSU (Biological study, unclassified); PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(pyruvate kinase allosteric inhibitors for therapeutic use)

RN 289060-07-7 CAPLUS

CN 1,3-Benzenedicarboxylic acid, 5-[4-(methoxycarbonyl)-2-[[[3-(trifluoromethyl)phenyl]amino]carbonyl]amino]phenoxy]-, dimethyl ester
(9CI) (CA INDEX NAME)



L6 ANSWER 2 OF 6 CAPLUS COPYRIGHT 2002 ACS

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* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

AB Title compds. [I; X = O, S; A = 1,4-C6H4, 1,3-C6H4, 1,7-naphthyl; L = H, 2,6-(CH3)2, 2-(CH3)3C, 6-(CH3)3C; R1 = CH3NHCO, CH3CH2NHCO, (CH3)3CNHCO, CH3(CH2)5NHCO, CF3NHCO, C6H5NHCO, 2-CH3C6H4NHCO, 3-CH3C6H4NHCO, 4-CH3C6H4NHCO, 2,6-(CH3)2C6H3NHCO, 4-CF3C6H4NHCO, 2,3-F2C6H4NHCO; q = 0-8; m = 0-8; n = 0-8] and pharmacol. acceptable salts, which are useful as therapeutic and/or preventive agents for diabetes, hyperlipemia, arteriosclerosis, **cancers**, are prepd. Thus, the title compd. II was prepd.

AN 2000:742094 CAPLUS

DN 133:296435

TI Preparation of amine derivatives useful agents for diabetes, hyperlipemia, arteriosclerosis, and **cancer**

IN Fujita, Takashi; Wada, Kunio; Oguchi, Minoru; Honma, Hidehito; Fujiwara, Toshihiko

PA Sankyo Company, Limited, Japan

SO PCT Int. Appl., 208 pp.

CODEN: PIXXD2

DT Patent

LA Japanese

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	WO 2000061581	A1	20001019	WO 2000-JP2216	20000406
	W: AU, BR, CA, CN, CZ, HU, ID, IL, IN, KR, MX, NO, NZ, PL, RU, TR, US, ZA				
	RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE				
	JP 2000351779	A2	20001219	JP 2000-104702	20000406
	EP 1167366	A1	20020102	EP 2000-915362	20000406
	R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, FI				
	BR 2000009594	A	20020604	BR 2000-9594	20000406
	NO 2001004847	A	20011207	NO 2001-4847	20011005
PRAI	JP 1999-99981	A	19990407		
	WO 2000-JP2216	W	20000406		

OS MARPAT 133:296435

IT 301548-73-2P

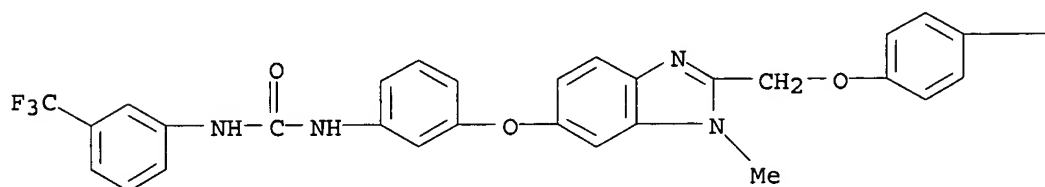
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

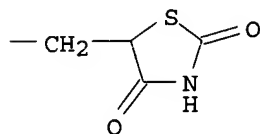
(prepn. of amine derivs. as useful agents for diabetes, hyperlipemia, arteriosclerosis, and **cancer**)

RN 301548-73-2 CAPLUS

CN Urea, N-[3-[[2-[[4-[(2,4-dioxo-5-thiazolidinyl)methyl]phenoxy)methyl]-1-methyl-1H-benzimidazol-6-yl]oxy]phenyl]-N'-[3-(trifluoromethyl)phenyl]-(9CI) (CA INDEX NAME)

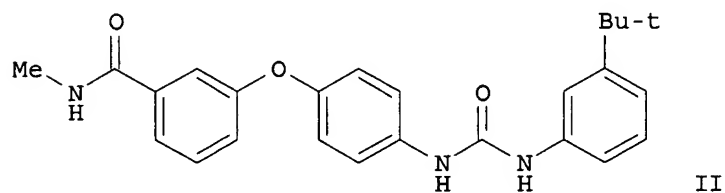
PAGE 1-A





RE.CNT 16 THERE ARE 16 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 3 OF 6 CAPLUS COPYRIGHT 2002 ACS
GI



L6 ANSWER 6 OF 6 CAPLUS COPYRIGHT 2002 ACS

AB ET-18-O-CH₃ (1-O-octadecyl-2-O-methyl-sn-glycero-3-phosphocholine) is an antiproliferative agent, blocking the growth of **cancer** cells both in vitro and in vivo. However, there is controversy regarding the mechanism leading to its antiproliferative effects. CoA-independent transacylase (CoA-IT) is an enzyme that remodels arachidonate between specific phospholipid donor and acceptor mols. in a variety of mammalian cells. ET-18-O-CH₃ was a potent inhibitor of CoA-IT (IC₅₀, 0.5 .mu.M), and kinetic anal. revealed that its inhibition was competitive with the lyso-phospholipid substrate. The goal of the current study was to explore the connection between inhibition of CoA-IT and antiproliferative effects using several structurally distinct inhibitors of CoA-IT. ET-18-O-CH₃ and other inhibitors of CoA-IT were found to inhibit cell proliferation and thymidine incorporation into the DNA, as well as to induce apoptosis in human HL-60 monocytic leukemia cells. The mechanism of apoptosis induced by ET-18-O-CH₃ appeared to be different from that induced by tumor necrosis factor; the former failed to activate NF-.kappa.B, whereas tumor necrosis factor did. Closer examn. of the pharmacol. of apoptosis in this model revealed that compds. that were structurally related to CoA-IT inhibitors, but lacked CoA-IT inhibitory activity, also failed to induce apoptosis. In addn., compds. that inhibited other enzymes that participate in arachidonic acid metab., cyclooxygenase, 5-lipoxygenase and phospholipase A₂, did not induce apoptosis. Taken together, these results demonstrate that inhibition of CoA-IT can be linked to blockade of proliferation and the induction of apoptosis in HL-60 cells.

AN 1996:702444 CAPLUS

DN 126:166148

TI Inhibitors of coenzyme A-independent transacylase induce apoptosis in human HL-60 cells

AU Winkler, James D.; Eris, Tamer; Sung, Chiu-Mei; Chabot-Fletcher, Marie; Mayer, Ruth J.; Surette, Marc E.; Chilton, Floyd H.

CS Dep. Immunopharmacol. Med. Chem., SmithKline Beecham Pharmaceuticals, King of Prussia, PA, USA

SO Journal of Pharmacology and Experimental Therapeutics (1996), 279(2), 956-966

CODEN: JPETAB; ISSN: 0022-3565

PB Williams & Wilkins

DT Journal

LA English

IT 162793-63-7, Skf 45905

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); BIOL (Biological study)

(inhibitors of CoA-independent transacylase induce apoptosis in human HL-60 cells)

RN 162793-63-7 CAPLUS

CN Benzenesulfonic acid, 4,5-dichloro-2-[2-[[[4-chloro-3-(trifluoromethyl)phenyl]amino]carbonyl]amino]-4-(trifluoromethyl)phenoxy]-(9CI) (CA INDEX NAME)

